- 5. Section 508 Compliance-Contestants must acknowledge that they understand that, as a pre-requisite to any subsequent acquisition by FAR contract or other method, they may be required to make their proposed solution compliant with Section 508 accessibility and usability requirements at their own expense. Any electronic information technology that is ultimately obtained by HHS for its use, development, or maintenance must meet Section 508 accessibility and usability standards. Past experience has demonstrated that it can be costly for solution-providers to "retrofit" solutions if remediation is later needed. The HHS Section 508 Evaluation Product Assessment Template, available at http://www.hhs.gov/web/508/ contracting/technology/vendors.html, provides a useful roadmap for developers to review. It is a simple, web-based checklist utilized by HHS officials to allow vendors to document how their products do or do not meet the various Section 508 requirements.
- 6. Functionality/Accuracy—A Submission may be disqualified if the application fails to function as expressed in the description provided by the user, or if the application provides inaccurate or incomplete information.
- 7. Security—Submissions must be free of malware. Contestant agrees that ONC may conduct testing on the application to determine whether malware or other security threats may be present. ONC may disqualify the application if, in ONC's judgment, the application may damage government or others' equipment or operating environment.

Additional Information

Ownership of intellectual property is determined by the following:

- Patient Application category entrants retain title and full ownership in and to their submission. Entrants expressly reserve all intellectual property rights not expressly granted under the challenge agreement.
- Developer Tools category entrants are required to post their tools on GitHub to be made available via open source.
- By participating in the challenge, each entrant hereby irrevocably grants to Sponsor and Administrator a limited, non-exclusive, royalty-free, worldwide license and right to reproduce, publically perform, publically display, and use the Submission to the extent necessary to administer the challenge, and to publically perform and publically display the Submission, including, without limitation, for

advertising and promotional purposes relating to the challenge.

Authority: 15 U.S.C. 3719

Dated: May 28, 2013. Farzad Mostashari,

National Coordinator for Health Information Technology.

[FR Doc. 2013–13128 Filed 6–3–13; 8:45 am]

BILLING CODE 4150-45-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Assistant Secretary for Planning and Evaluation; Advisory Council on Alzheimer's Research, Care, and Services

AGENCY: Office of the Assistant Secretary for Planning and Evaluation, Department of Health and Human Services.

ACTION: Request for Nominations.

SUMMARY: HHS is soliciting nominations for a new, non-Federal member of the Advisory Council on Alzheimer's Research, Care, and Services. Specifically, the position is for someone with a diagnosis of Alzheimer's disease or a related dementia. Nominations should include the nominee's contact information (current mailing address, email address, and telephone number) and current curriculum vitae or resume. Nominations submitted within the past 6 months for other positions on the Advisory Council on Alzheimer's Research, Care, and Services will be considered for this position.

DATES: Submit nominations by email or FedEx or UPS before COB on June 14, 2013.

ADDRESSES: Nominations should be sent to Helen Lamont at helen.lamont@hhs.gov; Helen Lamont, Ph.D., Office of the Assistant Secretary for Planning and Evaluation, Room 424E Humphrey Building, Department of Health and Human Services, 200 Independence Avenue SW., Washington, DC 20201.

FOR FURTHER INFORMATION CONTACT: Helen Lamont (202) 690–7996, helen.lamont@hhs.gov.

SUPPLEMENTARY INFORMATION: The Advisory Council on Alzheimer's Research, Care, and Services meets quarterly to discuss programs that impact people with Alzheimer's disease and related dementias and their caregivers. The Advisory Council makes recommendations about ways to reduce the financial impact of Alzheimer's disease and related dementias and to improve the health outcomes of people with these conditions. The Advisory

Council provides feedback on the National Plan to Address Alzheimer's Disease. On an annual basis, the Advisory Council shall evaluate the implementation of the recommendations through an updated national plan.

The Advisory Council consists of designees from Federal agencies including the Centers for Disease Control and Prevention, Administration on Aging, Centers for Medicare and Medicaid Services, Indian Health Service, Office of the Director of the National Institutes of Health, National Science Foundation, Department of Veterans Affairs, Food and Drug Administration, Agency for Healthcare Research and Quality, and the Surgeon General. The Advisory Council also consists of 13 non-federal members selected by the Secretary who are Alzheimer's patient advocates (2), Alzheimer's caregivers (2), health care providers (2), representatives of State health departments (2), researchers with Alzheimer's-related expertise in basic, translational, clinical, or drug development science (2), voluntary health association representatives (2), and a person with a diagnosis of Alzheimer's disease or a related dementia. Members serve as Special Government Employees. This announcement is seeking nominations for a person with a diagnosis of Alzheimer's disease or a related dementia who is not a Federal employee. This person will serve a twoyear term.

Dated: May 28, 2013.

Donald B. Moulds,

Acting Assistant Secretary for Planning and Evaluation.

[FR Doc. 2013–13127 Filed 6–3–13; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [Docket No. FDA-2013-N-0271]

Availability of Masked and Deidentified Non-Summary Safety and Efficacy Data; Request for Comments

ACTION: Notice; request for comments.

AGENCY: Food and Drug Administration,

SUMMARY: The Food and Drug Administration (FDA) is seeking public comments from interested persons on the proposed availability of deidentified and masked data derived from medical product applications. Improving the efficiency and