

to residents of Illinois with funding from the Retirement Research Foundation. Additional funds are needed to leverage the foundation's funding, in order to ensure that the current provision of services to Illinois residents will be continued. This supplementary funding would be provided for the approved period.

Authority: This program is authorized under Title II of the Older Americans Act (OAA) (42 U.S.C. 3032), as amended by the Older Americans Act Amendments of 2006, Public Law 109–365.

(Catalog of Federal Domestic Assistance 93.048).

Dated: August 14, 2015.

Kathy Greenlee,

Assistant Secretary for Aging and Administrator, Administration for Community Living.

[FR Doc. 2015–22630 Filed 9–8–15; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Community Living

Announcement of the Intent To Award Single-Source Grants to the National Association of Area Agencies on Aging and the National Association of States United for Aging and Disabilities

AGENCY: Administration for Community Living, HHS.

ACTION: Notice.

SUMMARY: The Administration for Community Living (ACL) announces the intent to award, subject to the availability of funds, single-source grant awards in the amount of \$335,000 to the National Association of Area Agencies on Aging (n4a) and \$153,500 to the National Association of State United for Aging and Disabilities (NASUAD). The awards will continue supporting and stimulating the ongoing work by these organizations to further develop and assist states and community-based organizations with building their business capacity for managed long-term services and supports and delivery system reform. CFDA Numbers: 93.048

DATES: The awards will be issued for a project period of September 30, 2015 through September 29, 2016.

FOR FURTHER INFORMATION CONTACT: Marisa Scala-Foley, Office of Integrated Care Innovations, Administration for Community Living, 1 Massachusetts Avenue NW., Washington, DC 20001. Telephone: 202–357–3516; Email: Marisa.Scala-Foley@acl.hhs.gov.

SUPPLEMENTARY INFORMATION: In 2012, n4a and NASUAD were awarded grants from ACL to build the business capacity of state and community-based aging and disability organizations for managed long-term services and supports (MLTSS). This one-year of grant funding, through continuation grant, will continue to support n4a and NASUAD in their efforts to:

- Identify and track emerging trends, best practices, barriers, lessons learned and progress in the aging and disability networks' integration into MLTSS and delivery system reform;
- increase state and community-based aging and disability organizations' capacity, readiness and involvement in the provision of MLTSS through the provision of broad-based and targeted technical assistance, education and training; and
- develop products that complement and enhance the first two areas of focus.

This program is authorized under the Older Americans Act of 1965, as amended in 2006, Public Law 109–365.

Dated: August 28, 2015.

Sharon Lewis,

Principal Deputy Administrator, Administration for Community Living.

[FR Doc. 2015–22631 Filed 9–8–15; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2013–D–1213]

Use of Donor Screening Tests To Test Donors of Human Cells, Tissues, and Cellular and Tissue-Based Products for Infection With *Treponema pallidum* (Syphilis); Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of a document entitled “Use of Donor Screening Tests to Test Donors of Human Cells, Tissues and Cellular and Tissue-Based Products for Infection with *Treponema pallidum* (Syphilis); Guidance for Industry.” The guidance document provides establishments that make donor eligibility determinations for donors of human cells, tissues, and cellular and tissue-based products (HCT/P Establishments) with updated recommendations concerning donor testing for evidence of *Treponema pallidum* (*T. pallidum*) infection, the

etiologic agent of syphilis. HCT/P Establishments must, as required under Federal regulations, test a donor specimen for evidence of *T. pallidum* infection using appropriate FDA-licensed, approved, or cleared donor screening tests, in accordance with the manufacturer's instructions, unless an exception to this requirement applies. The guidance clarifies that FDA does not consider diagnostic tests or pre-amendment devices (which have not been licensed, approved, or cleared) to be adequate for use in donor testing for *T. pallidum* infection under the criteria specified in Federal regulations. The guidance announced in this notice finalizes the draft guidance of the same title, dated October 2013. The recommendations in the guidance announced in this notice supersedes those recommendations for testing HCT/P donors for evidence of *T. pallidum* infection contained in the document entitled “Guidance for Industry: Eligibility Determination for Donors of Human Cells, Tissues, and Cellular and Tissue-Based Products (HCT/Ps),” dated August 2007.

DATES: Submit either electronic or written comments on Agency guidances at any time.

ADDRESSES: Submit written requests for single copies of the guidance to the Office of Communication, Outreach and Development, Center for Biologics Evaluation and Research (CBER), Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 71, Rm. 3128, Silver Spring, MD 20993–0002. Send one self-addressed adhesive label to assist the office in processing your requests. The guidance may also be obtained by mail by calling CBER at 1–800–835–4709 or 240–402–8010. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the guidance document.

Submit electronic comments on the guidance to <http://www.regulations.gov>. Submit written comments to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT: Paul E. Levine, Jr., Center for Biologics Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 71, Rm. 7301, Silver Spring, MD 20993–0002, 240–402–7911.

SUPPLEMENTARY INFORMATION:

I. Background

FDA is announcing the availability of a document entitled “Use of Donor Screening Tests to Test Donors of