In addition, CERCLA provides ATSDR with the authority to prepare toxicological profiles for substances not found on the SPL. CERCLA authorizes ATSDR to establish and maintain a inventory of literature, research, and studies on the health effects of toxic substances (CERCLA Section 104(i)(1)(B)); to respond to requests for health consultations (CERCLA Section 104(i)(4)); and to support the site-specific response actions conducted by the agency.

Availability


Pamela J. Protzel Berman, Director, Office of Policy, Partnerships and Planning, Agency for Toxic Substances and Disease Registry.

FOR FURTHER INFORMATION CONTACT: contact Melvenia Wright, U.S. Department of Health and Human Services, Administration on Disability Rights Network (NDRN) 90HAVA0001 and the National Federation of the Blind (NFB) 90HAVA0002.

SUMMARY: The Administration for Community Living (ACL) announces the intent to award an administrative supplement to the current Help America Vote Act (HAVA) Training and Technical Assistance (T/TA) grantees held by the National Disability Rights Network (NDRN) 90HAVA0001 and the National Federation of the Blind (NFB) 90HAVA0002.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Community Living

Announce the Intent To Award an Administrative Supplement

ACTION: Announcing the intent to award an Administrative Supplement to two (2) Help America Vote Act (HAVA) Training and Technical Assistance (T/TA) grantees held by the National Disability Rights Network (NDRN) 90HAVA0001 and the National Federation of the Blind (NFB) 90HAVA0002.

SUMMARY: The Administration for Community Living (ACL) announces the intent to award an administrative supplement to the current Help America Vote Act (HAVA) Training and Technical Assistance (T/TA) grantees held by the National Disability Rights Network (NDRN) 90HAVA0001 and the National Federation of the Blind (NFB). The purpose of the HAVA programs are designed to establish and improve participation in the election process for individuals with a full range of disabilities. In each eligible state and territory, seven percent of HAVA funds are set aside for the Protection and Advocacy Systems (P&As) to ensure that individuals with disabilities have the opportunity to participate in every step of the voting process. After receiving training and technical assistance, P&As may train others on the availability of accessible voting equipment and its use. The administrative supplement for FY 2018 will be in the amount of $122,721 bringing the total award for FY 2018 to $462,590.

Program Name: Help America Vote Act Training and Technical Assistance.

Recipients: National Disability Rights Network (NDRN) and National Federation of the Blind (NFB).

Period of Performance: The supplement award will be issued for the second year of the two-year project period of September 1, 2018, through August 30, 2019.


Award Type: Administrative Supplement.


Basis for Award: The additional funding will not be used to begin new projects. The funding will be used to increase NDRN’s capacity building efforts to provide training and technical assistance to the Protection and Advocacy Systems in the electoral process and NFB will be able to attend voting related conferences, conduct voting outreach campaigns and translate materials into Spanish.

FOR FURTHER INFORMATION CONTACT: For further information or comments regarding this program supplement, contact Melvenia Wright, U.S. Department of Health and Human Services, Administration on Disabilities, Administration on Intellectual and Developmental Disabilities: telephone (202) 795–7472; email Melvenia.Wright@acl.hhs.gov.

Dated: July 26, 2018.

Lance Robertson, Administrator and Assistant Secretary for Aging.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Revocation of Authorization of Emergency Use of an In Vitro Diagnostic Device for Detection of Ebola Virus

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the revocation of the Emergency Use Authorization (EUA) (the Authorization) issued to Zalgen Labs, LLC for the ReEBOV Antigen Rapid Test. FDA revoked this Authorization on May 18, 2018, under the Federal Food, Drug, and Cosmetic Act (FD&C Act), as requested by Zalgen Labs, LLC by letter dated March 1, 2018. The revocation, which includes an explanation of the reasons for revocation, is reprinted in this document.

DATES: The Authorization is revoked as of May 18, 2018.

ADDRESSES: Submit written requests for single copies of the revocation to the Office of Counterterrorism and Emerging Threats, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 1, Rm. 4336, Silver Spring, MD 20993–0002. Send one self-addressed adhesive label to assist that office in processing your request or include a Fax number to which the documentation may be sent. See the SUPPLEMENTARY INFORMATION section for electronic access to the revocation.

FOR FURTHER INFORMATION CONTACT: Michael Maier, Office of Counterterrorism and Emerging Threats, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 1, Rm. 4336, Silver Spring, MD 20993–0002, 301–796–8510 (this is not a toll-free number).

SUPPLEMENTARY INFORMATION:

I. Background

Section 564 of the FD&C Act (21 U.S.C. 360bbb–3) as amended by the Project BioShield Act of 2004 (Pub. L. 108–276) and the Pandemic and All-Hazards Preparedness Reauthorization Act of 2013 (Pub. L. 113–5) allows FDA to strengthen the public health protections against biological, chemical, nuclear, and radiological agents. Among other things, section 564 of the FD&C Act allows FDA to authorize the use of an unapproved medical product for an unapproved use of an approved medical product in certain situations. On February 24, 2015, FDA issued an EUA to Corgenix, Inc. for the ReEBOV Antigen Rapid Test, subject to the terms of the Authorization. Notice of the issuance of the Authorization was published in the Federal Register on June 5, 2015 (80 FR 32140), as required by section 564(h)(1) of the FD&C Act. In response to requests from Zalgen Labs, LLC and Corgenix, Inc. to transfer ownership of the EUA for the ReEBOV Antigen Rapid Test, Corgenix, Inc. to Zalgen Labs, LLC, FDA amended and reissued the EUA to Zalgen Labs, LLC.

II.PyObject Method

Summary: The Food and Drug Administration (FDA) is announcing the revocation of the Emergency Use Authorization (EUA) (the Authorization) issued to Zalgen Labs, LLC for the ReEBOV Antigen Rapid Test. FDA revoked this Authorization on May 18, 2018, under the Federal Food, Drug, and Cosmetic Act (FD&C Act), as requested by Zalgen Labs, LLC by letter dated March 1, 2018. The revocation, which includes an explanation of the reasons for revocation, is reprinted in this document.

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