proposed collection of information; (c) the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Dated: August 8, 2011.

#### Steven M. Hanmer,

Reports Clearance Officer.

[FR Doc. 2011–20495 Filed 8–11–11; 8:45 am]

BILLING CODE 4184-09-M

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Administration for Children and Families

# Announcement of a Single Source Grant Award to the Tribal Law and Policy Institute

**AGENCY:** Children's Bureau, Administration on Children, Youth and Families, HHS.

**ACTION:** Notice to award a single source program expansion supplement grant to the Tribal Law and Policy Institute, located in West Hollywood, CA, to support activities of the National Resource Center for Tribes under the Tribal Maternal, Infant, Early Childhood Home Visiting Program.

CFDA Number: 93.508.

**Statutory Authority:** Social Security Act, Title V, Section 511 (42 U.S.C. 701), as amended by the Patient Protection and Affordable Care Act of 2010 (ACA), Pub. L. 111–148.

**SUMMARY:** The Administration for Children and Families (ACF), Administration on Children, Youth and Families (ACYF), Children's Bureau (CB) announces the award of a single source program expansion supplement grant to the Tribal Law and Policy Institute, West Hollywood, CA, for the National Resource Center (NRC) for Tribes. The program expansion supplement funds will be used to provide technical assistance and support for the planning, development and implementation of the Tribal Maternal, Infant and Early Childhood Home Visiting program.

The NRC for Tribes will provide technical assistance to ACF Tribal Home Visiting grantees to enhance their capacity to plan for and implement high-quality, evidence-based, and evidence-informed programs.

Implementation of the NRC4Tribes work

will include engaging, assessing, informing and supporting culturally-appropriate Tribal home visiting services that are part of coordinated early childhood systems in the American Indian and Alaska Natives (AIAN) communities and that support quality and effectiveness of services for AIAN children, youth, and families, which leads to increased safety, permanency, and well-being for children.

The Tribal Law and Policy Institute NRC for Tribes and its partner agencies are uniquely qualified to provide training and technical assistance to Tribes based upon their experience, expertise, and commitment to increasing cultural competency and sensitivity to the Tribal point of view in training and technical assistance. The NRC for Tribes expertise in Tribal culture, child maltreatment prevention, collaboration, evaluation, and implementation of evidence-based programs and practices makes them an appropriate recipient of supplemental funds to carry out this project.

Amount of Award: \$150,000. Project Period: May 15, 2011 to September 30, 2011.

### FOR FURTHER INFORMATION CONTACT:

Roshanda Shoulders, Children's Bureau, 1250 Maryland Ave., SW., 8th Floor, Washington, DC 20024. Telephone: (202) 401–5323. E-mail: roshanda.shoulders@acf.hhs.gov.

Dated: August 2, 2011.

### Bryan Samuels,

Commissioner, Administration on Children, Youth and Families.

[FR Doc. 2011–20278 Filed 8–11–11; 8:45 am]

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. FDA-2011-N-0271]

Harmful and Potentially Harmful Constituents in Tobacco Products and Tobacco Smoke; Request for Comments

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice; request for comments.

SUMMARY: The Food and Drug Administration (FDA) is requesting comments, including scientific and other information, concerning the harmful and potentially harmful constituents (HPHCs) in tobacco products and tobacco smoke. This information will assist the Agency in

establishing a list of HPHCs in tobacco products and tobacco smoke (the HPHC list).

**DATES:** Submit either electronic or written comments by October 11, 2011.

**ADDRESSES:** Submit electronic comments 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. Identify comments with the docket number found in brackets in the heading of this document.

#### FOR FURTHER INFORMATION CONTACT:

Carol Drew, Center for Tobacco Products, Food and Drug Administration, 9200 Corporate Blvd., Rockville, MD 20850–3229, 877–287– 1373.

### SUPPLEMENTARY INFORMATION:

#### I. Background

On June 22, 2009, the President signed the Family Smoking Prevention and Tobacco Control Act (Tobacco Control Act) (Pub. L. 111-31) into law. The Tobacco Control Act amended the Federal Food, Drug, and Cosmetic Act (the FD&C Act) (21 U.S.C. 301 et seq.) by, among other things, adding a new chapter granting FDA important new authority to regulate the manufacture, marketing, and distribution of tobacco products to protect the public health generally and to reduce tobacco use by minors. Section 904(e) of the FD&C Act (21 U.S.C. 387d(e)), as added by the Tobacco Control Act, requires FDA to establish, and periodically revise as appropriate, "a list of harmful and potentially harmful constituents, including smoke constituents, to health in each tobacco product by brand and by quantity in each brand and subbrand." Section 904(e) of the FD&C Act also requires that FDA "publish a public notice requesting the submission by interested persons of scientific and other information concerning the harmful and potentially harmful constituents in tobacco products and tobacco smoke.'

The Agency has solicited scientific and other information from interested persons and has developed a list of tobacco product constituents it currently believes are harmful or potentially harmful to health. Although the Agency's work to date reflects consideration of substantial scientific and other information, we believe that additional information from the public may be beneficial to the Agency before it establishes the list described in section 904(e) of the FD&C Act. We are therefore publishing the Agency's