

not be accepted as proof of timely mailing.)

2. Late Applications

Applications that do not meet the criteria in 1.A. or 1.B. above are considered late applications. Late applications will not be considered in the current competition and will be returned to the applicant.

Where To Obtain Additional Information

A complete program description, information on application procedures, an application package, and business management technical assistance may be obtained from Van Malone, Grants Management Officer, Grants Management Branch, Procurement and Grants Office, Centers for Disease Control and Prevention (CDC), 255 East Paces Ferry Road, NE., Room 300, Mail Stop E-15, Atlanta, Georgia 30305, telephone (404) 842-6575, Email address vxm7@cdc.gov. The announcement will be available on one of two Internet sites on the publication date: CDC's home page at <http://www.cdc.gov>, or at the Government Printing Office home page (including free access to the **Federal Register**) at <http://www.access.gpo.gov>.

Programmatic technical assistance may be obtained from Dr. Richard Steketee or Dr. Marta Gwinn, Division of HIV/AIDS Prevention, National Center for HIV, STD, TB Prevention, Centers for Disease Control and Prevention (CDC), 1600 Clifton Road, NE., Mail Stop E-46, Atlanta, Georgia 30333, telephone (404) 639-2090. Eligible applicants are encouraged to call before developing and submitting their application. Please refer to Announcement Number 797 when requesting information.

Potential applicants may obtain a copy of "Healthy People 2000" (Full Report: Stock No. 017-001-00474-0) or "Healthy People 2000" (Summary Report: Stock No. 017-001-00473-1) referenced in the Introduction from the Superintendent of Documents, Government Printing Office, Washington, DC 20402-9325, telephone (202) 512-1800.

Dated: July 3, 1997.

Joseph R. Carter,

Acting Associate Director for Management and Operations, Centers for Disease Control and Prevention (CDC).

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Disease, Disability, and Injury Prevention and Control Special Emphasis Panel (SEP): Cooperative Agreement for a National Center for the Prevention of Childhood Agricultural Injury, Program Announcement 737: Meeting

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92-463), the Centers for Disease Control and Prevention (CDC) announces the following committee meeting.

Name: Disease, Disability, and Injury Prevention and Control SEP: Cooperative Agreement for a National Center for the Prevention of Childhood Agricultural Injury, Program Announcement 737.

Time and Date: 8:30 a.m.-4:30 p.m., August 4, 1997.

Place: Corporate Square Building 11, Conference Room A, Corporate Square Boulevard, Atlanta, Georgia 30326.

Status: Closed.

Matters to be Discussed: The meeting will include the review, discussion, and evaluation of applications received in response to Program Announcement 737.

The meeting will be closed to the public in accordance with provisions set forth in section 552b(c) (4) and (6), Title 5 U.S.C., and the Determination of the Associate Director for Management and Operations, CDC, pursuant to Pub. L. 92-463.

Contact Person for More Information: Ann Cronin, Office of Extramural Coordination and Special Projects, National Institute for Occupational Safety and Health, CDC, M/S D36, 1600 Clifton Road, NE, Atlanta, Georgia 30333, telephone 404/639-2277.

Dated: July 03, 1997.

Carolyn J. Russell,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention (CDC).

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 97D-0261]

Frequently Asked Questions About the New FDA Tobacco Regulations: Draft Guidance; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration is announcing the

availability of a draft guidance entitled "Frequently Asked Questions About the New FDA Tobacco Regulations." The draft guidance is intended to address the questions most frequently asked by retailers, consumers, and others about the age and identification requirements of the final rule restricting the sale of cigarettes and smokeless tobacco to protect children and adolescents.

DATES: Submit written comments on the draft guidance by September 8, 1997.

ADDRESSES: The draft guidance entitled "Frequently Asked Questions About the New FDA Tobacco Regulations," is available on the Internet at <http://www.fda.gov/>, or a paper copy may be ordered free of charge by calling 1-888-FDA-4KIDS.

Submit written comments on the draft guidance to the Dockets Management Branch (HFA-305), Food and Drug Administration, 12420 Parklawn Dr., rm. 1-23, Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT: Anne M. Kirchner, Office of Policy (HF-11), Food and Drug Administration, 5600 Fishers Lane, rm. 14-72, Rockville, MD 20857, 301-827-0867.

SUPPLEMENTARY INFORMATION: In the **Federal Register** of August 28, 1996 (61 FR 44396), FDA issued a final rule to restrict the sale and distribution of cigarettes and smokeless tobacco in order to protect children and adolescents (21 CFR part 897). The final rule covers three general classes of nicotine-containing tobacco products: Cigarettes, loose cigarette tobacco, and smokeless tobacco. The final rule applies to manufacturers, distributors, retailers, and importers who make, distribute, sell, and import such products.

Since February 28, 1997, the final rule has prohibited retailers from selling cigarettes, loose cigarette tobacco, or smokeless tobacco to persons under the age of 18, and has required retailers to verify the age of customers under the age of 27 by checking an identification (ID) card which contains the bearer's photograph and birth date.

Before the age and ID requirements took effect, FDA officials held a series of public meetings in 10 metropolitan areas and produced a national videoconference to explain the new requirements and to answer questions from retailers, consumers, public health officials, and others. FDA agreed to make available written answers to the questions most frequently asked at these meetings.

The draft guidance that FDA is making available answers these questions, as well as questions that FDA has received on its toll-free hotline and