

**SUPPLEMENTARY INFORMATION:** Pursuant to Section 6(f) of the Federal Trade Commission Act, 38 Stat. 721, 15 U.S.C. 46 and Section 2.34 of the Commission's Rules of Practice (16 CFR 2.34), notice is hereby given that the above-captioned consent agreement containing a consent order to cease and desist, having been filed with and accepted by the Commission, has been placed on the public record for a period of thirty (30) days. The following Analysis to Aid Public Comment describes the terms of the consent agreement, and the allegations in the complaint. An electronic copy of the full text of the consent agreement package can be obtained from the FTC Home Page (for October 11, 2001), on the World Wide Web, at "<http://www.ftc.gov/os/2001/10index.htm>." A paper copy can be obtained from the FTC Public Reference Room, Room H-130, 600 Pennsylvania Avenue, NW., Washington, DC 20580, either in person or by calling (202) 326-3627.

Public comment is invited. Comments should be directed to: FTC/Office of the Secretary, Room 159, 600 Pennsylvania Ave., NW., Washington, DC 20580. Two paper copies of each comment should be filed, and should be accompanied, if possible, by a 3½ inch diskette containing an electronic copy of the comment. Such comments or views will be considered by the Commission and will be available for inspection and copying at its principal office in accordance with Section 4.9(b)(6)(ii) of the Commission's Rules of Practice (16 CFR 4.9(b)(6)(ii)).

#### **Analysis of Proposed Consent Order To Aid Public Comment**

The Federal Trade Commission has accepted, subject to final approval, an agreement to a proposed consent order from respondent FanBuzz, Inc.

The proposed consent order has been placed on the public record for thirty (30) days for reception of comments by interested persons. Comments received during this period will become part of the public record. After thirty (30) days, the Commission will again review the agreement and the comments received and will decide whether it should withdraw from the agreement and take other appropriate action or make final the agreement's proposed order.

This matter concerns practices related to the sale of textile products by means of an Internet catalog. The Commission's complain charges that respondent violated the Federal Trade Commission Act, 15 U.S.C. 41 *et seq.*, and the Textile Fiber Products Identification Act, 15 U.S.C. 70 *et seq.*,

by failing to disclose in its Internet catalog whether products offered for sale were made in the United States, imported, or both.

Part I of the proposed consent order prohibits future violations of the Textile Fiber Products Identification Act and Commission rules and regulations, found at 16 CFR Part 303, implementing the requirements of the statute.

Part II of the proposed order requires the respondent, for five years after the date of issuance of the Order, to maintain records demonstrating compliance with the Order, including: (a) Copies of mail order catalogs and mail order promotional materials, as defined in 16 CFR 303.1(u), that offer textile products for direct sale to consumers; and (b) complaints and other communications with consumers, government agencies, or consumer protection organizations, pertaining to country-of-origin disclosures for textile products.

Part III of the proposed order requires the respondent to distribute copies of the order to certain company officials and employees. Part IV of the proposed order requires the respondent to notify the Commission of any change in the corporation that may affect compliance obligations under the order. Part V of the proposed order requires the respondent to file one or more compliance reports. Part VI of the proposed order is a provision whereby the order, absent certain circumstances, terminates twenty years from the date of issuance.

The purpose of this analysis is to facilitate public comments on the proposed consent order. It is not intended to constitute an official interpretation of the agreement and proposed order or to modify in any way their terms.

By direction of the Commission.

**Donald S. Clark,**

*Secretary.*

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## **DEPARTMENT OF HEALTH AND HUMAN SERVICES**

### **Centers for Disease Control and Prevention**

**[30Day-55-01]**

#### **Agency Forms Undergoing Paperwork Reduction Act Review**

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under

review by the Office of Management and Budget (OMB) in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 639-7090. Send written comments to CDC, Desk Officer, Human Resources and Housing Branch, New Executive Office Building, Room 10235, Washington, DC 20503. Written comments should be received within 30 days of this notice.

#### **Proposed Project**

Pulmonary Function Testing Course Approval Program, 29 CFR 1910.1043 (OMB No. 0920-0138)—EXTENSION—The National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC). The mission of the National Institute for Occupational Safety and Health is to promote safety and health at work for all people through research and prevention.

NIOSH has responsibility under the Cotton Dust Standard, 29 CFR 1910.1043, for approving courses to train technicians to perform pulmonary function testing in the Cotton Dust Industry. Successful completion of a NIOSH-approved course is mandatory under the Standard. To carry out its responsibility, NIOSH maintains a Pulmonary Function Testing Course Approval Program. The program consists of an application submitted by potential sponsors who seek NIOSH approval to conduct courses, and if approved, notification to NIOSH of any course or faculty changes during the period of approval. The application form and addend materials, including agenda, vitae and course materials, is reviewed by the National Institute for Occupational Safety and Health to determine if the applicant has developed a program which adheres to the criteria required in the Standard. Following approval, any subsequent changes to the course are submitted by course sponsors via letter and reviewed by NIOSH staff to assure that changes in faculty or course content continue to meet course requirements. Applications and materials to be a course sponsor and carry out training are submitted voluntarily by institutions and organizations from throughout the country. This is required for NIOSH to evaluate a course to determine whether it meets the criteria in the Standard and whether technicians will be adequately trained as mandated under the Standard. The estimated annual burden hours for this data collection is 66 hours.

Respondents	Number of respondents	Number of responses	Avg. burden/response (in hrs.)
Initial Application .....	5	1	3.5
Annual Letter .....	53	1	45/60
Report of Course Changes .....	12	1	45/60

Dated: October 10, 2001.

**Nancy E. Cheal,**

*Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention.*

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Administration for Children and Families

#### Statement of Organization, Functions, and Delegations of Authority

This notice amends Part K of the Statement of Organization, Functions, and Delegations of Authority of the Department of Health and Human Services (DHHS), Administration for Children and Families (ACF) as follows: Chapter KA, Office of the Assistant Secretary for Children and Families (OAS) as last amended January 2, 1998 (63 FR 81-87) and Chapter KP, Office of the Deputy Assistant Secretary for Administration (ODASA) as last amended February 27, 2001 (66 FR 12525-28) and April 9, 2001 (66 FR 18487). This notice realigns the Executive Secretariat Office from the Office of the Deputy Assistant Secretary for Administration to the Office of the Assistant Secretary for Children and Families.

These Chapters are amended as follows:

I. Chapter KA, Office of the Assistant Secretary for Children and Families.

A. Delete KA.10 Organization in its entirety and replace with the following:

KA.10 Organization. The Office of the Assistant Secretary for Children and Families is headed by the Assistant Secretary who reports directly to the Secretary and consists of:

—The Office of the Assistant Secretary (KA).

—President's Committee on Mental Retardation Staff (KAD).

—The Executive Secretariat Office (KAF).

B. Amend KA.20 Functions to add the following new paragraph:

C. The Executive Secretariat Office (ExecSec) ensures that issues requiring the attention of the Assistant Secretary, Deputy Assistant Secretaries and/or

executive staff are addressed on a timely and coordinated basis and facilitates decisions on matters requiring immediate action including White House, Congressional and Secretarial assignments. The Office serves as the ACF liaison with the HHS Executive Secretariat. It receives, assesses and controls incoming correspondence and assignments to the appropriate ACF component(s) for response and action and provides assistance and advice to ACF staff on the development of responses to correspondence. The Office provides assistance to ACF staff on the use of the controlled correspondence system. The Office coordinates and/or prepares congressional correspondence; and tracks development of periodic reports and facilitates departmental clearances. The Director of the Executive Secretariat Office serves as the Freedom of Information Act Officer for ACF and coordinates hot line calls received by the Office of Inspector General and the General Accounting Office relating to ACF operations and personnel.

II. Chapter KP, Office of the Deputy Assistant Secretary for Administration.

A. Delete KP.00 Mission in its entirety and replace with the following:

KP.00 Mission. The Deputy Assistant Secretary for Administration serves as principal advisor and counsel to the Assistant Secretary for Children and Families on all aspects of personnel administration and management, information resource management, financial, grants policy and procurement issues, staff development and training activities, organizational development and organizational analysis, administrative services and facilities management and state systems policy. Oversees the ACF Equal Employment Opportunity and Civil Rights program and all special initiatives activities for ACF.

B. Amend KP.10 Organization to delete "Executive Secretariat Office (KPG)."

C. Amend KP.20 Functions to delete paragraph G, in its entirety.

Dated: September 28, 2001.

**Wade F. Horn,**

*Assistant Secretary for Children and Families.*

[FR Doc. 01-25995 Filed 10-15-01; 8:45 am]

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. 01N-0437]

#### Agency Information Collection Activities; Proposed Collection; Comment Request; New Animal Drugs for Investigational Use

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

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal agencies are required to publish notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice solicits comments on reporting and recordkeeping requirements for new animal drugs for investigational use.

**DATES:** Submit written or electronic comments on the collection of information by December 17, 2001.

**ADDRESSES:** Submit electronic comments on the collection of information to <http://www.accessdata.fda.gov/scripts/oc/dockets/edockethome.cfm>. Submit written comments on the collection of information to the Dockets Management Branch (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

**FOR FURTHER INFORMATION CONTACT:** Denver Presley, Office of Information Resources Management (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-1472.

**SUPPLEMENTARY INFORMATION:** Under the PRA (44 U.S.C. 3501-3520), Federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor.