

Toyota to retain documents relating to its compliance with the order. For most records, the order requires that the documents be retained for a five-year period. For the third party assessments and supporting documents, Franklin Toyota must retain the documents for a period of three years after the date that each assessment is prepared. Part VI requires dissemination of the order now and in the future to persons with responsibilities relating to the subject matter of the order. Part VII ensures notification to the FTC of changes in corporate status. Part VIII mandates that Franklin Toyota submit a compliance report to the FTC within 90 days, and periodically thereafter as requested. Part IX is a provision "sunsetting" the order after twenty (20) years, with certain exceptions.

The purpose of the analysis is to aid public comment on the proposed order. It is not intended to constitute an official interpretation of the proposed order or to modify its terms in any way.

By direction of the Commission.

Richard C. Donohue,
Acting Secretary.

[FR Doc. 2012-14372 Filed 6-12-12; 8:45 am]

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GENERAL SERVICES ADMINISTRATION

[FMR Bulletin-PBS-2012-03; Docket 2012-0002; Sequence 11]

Federal Management Regulation; FMR Bulletin PBS-2012-03; Redesignations of Federal Buildings

AGENCY: Public Buildings Service (PBS), General Services Administration (GSA).

ACTION: Notice of a bulletin.

SUMMARY: The attached bulletin announces the designation and redesignation of three Federal buildings.

Expiration Date: This bulletin announcement expires October 31, 2012. The building designation and redesignations remains in effect until canceled or superseded by another bulletin.

FOR FURTHER INFORMATION CONTACT: U.S. General Services Administration, Public Buildings Service (PBS), 1800 F Street NW., Washington, DC 20405, telephone number: (202) 501-1100.

Dated:

Dan Tangherlini,

Acting Administrator of General Services.

U.S. GENERAL SERVICES ADMINISTRATION

REDESIGNATIONS OF FEDERAL BUILDINGS

TO: Heads of Federal Agencies

SUBJECT: Redesignations of Federal Buildings

1. *What is the purpose of this bulletin?* This bulletin announces the designation and redesignation of three Federal buildings.

2. *When does this bulletin expire?* This bulletin announcement expires October 31, 2012. The building designation and redesignations remain in effect until canceled or superseded by another bulletin.

3. *Designation.* The name of the designated property (between the United States Federal Courthouse and the Ed Jones Building located at 109 South Highland Avenue in Jackson, Tennessee) is as follows:

M.D. Anderson Plaza
Jackson, TN 38301

4. *Redesignation.* The former and new names of the redesignated buildings are as follows:

Former name	New name
United States Courthouse, 80 Lafayette Street, Jefferson City, MO 65101.	Christopher S. Bond United States Courthouse, 80 Lafayette Street, Jefferson City, MO 65101.
United States Courthouse, 222 West 7th Avenue, Anchorage, AL 99501.	James M. Fitzgerald United States Courthouse, 222 West 7th Avenue, Anchorage, AL 99501.

5. *Who should we contact for further information regarding redesignation of these Federal buildings?* U.S. General Services Administration, Public Buildings Service (PBS), 1800 F Street, NW., Washington, DC 20405, telephone number: (202) 501-1100.

Dated: June 7, 2012

Dan Tangherlini,
Acting Administrator of General Services.

[FR Doc. 2012-14416 Filed 6-12-12; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Biennial Progress Report of the Interagency Coordinating Committee on the Validation of Alternative Methods (ICCVAM)

AGENCY: Division of the National Toxicology Program (DNTP), National Institute of Environmental Health

Sciences (NIEHS), National Institutes of Health (NIH).

ACTION: Availability of Report.

SUMMARY: The NTP Interagency Center for the Evaluation of Alternative Toxicological Methods (NICEATM) announces the availability of the *Biennial Progress Report 2010-2011: Interagency Coordinating Committee on the Validation of Alternative Methods*. The report was prepared in accordance with requirements of the ICCVAM Authorization Act of 2000 (42 U.S.C. 285l-3).

The *Biennial Progress Report* describes activities and progress by NICEATM and ICCVAM during the period from January 2010 through December 2011. During the past two years, NICEATM, ICCVAM, and ICCVAM member agencies contributed to the national and international endorsement and adoption of 14 new and updated alternative safety testing methods. Since ICCVAM was

established, NICEATM, ICCVAM, and the ICCVAM member agencies have contributed to the regulatory acceptance of over 50 alternative methods that can be used to protect the health of people, animals, and the environment while reducing, refining, and replacing animal use.

The *Biennial Progress Report* is available on the NICEATM-ICCVAM Web site at <http://iccvam.niehs.nih.gov/about/ICCVAMrpts.htm>. Copies can also be requested from NICEATM (see "ADDRESSES").

ADDRESSES: Requests for copies of the report should be sent by mail, fax, or email to Dr. William S. Stokes, Director, NICEATM, NIEHS, P.O. Box 12233, Mail Stop: K2-16, Research Triangle Park, NC 27709, (telephone) 919-541-2384, (fax) 919-541-0947, (email) niceatm@niehs.nih.gov. Courier address: NICEATM, NIEHS, Room 2034, 530 Davis Drive, Morrisville, NC 27560.

FOR FURTHER INFORMATION CONTACT: Dr. William S. Stokes, NICEATM Director (phone 919-541-2384 or niceatm@niehs.nih.gov).

SUPPLEMENTARY INFORMATION:

Background

The ICCVAM Authorization Act of 2000 established ICCVAM as a permanent interagency committee of NIEHS under NICEATM. The Act directs ICCVAM to coordinate interagency technical reviews of proposed new, revised, and alternative testing methods, including those that may reduce, refine (enhance animal well-being and lessen or avoid pain and distress), and replace animal use. ICCVAM prepares test method recommendations based on their scientific validity for regulatory safety testing, and submits these recommendations through the HHS Secretary (or designee) to U.S. Federal Agencies for adoption decisions.

A provision of the ICCVAM Authorization Act states that ICCVAM shall prepare "reports to be made available to the public on its progress under this Act," with the first report to be completed within 12 months of enactment of the Act, and subsequent reports to be made biennially thereafter. The fifth ICCVAM biennial progress report, which summarizes ICCVAM activities and accomplishments for the years 2010 and 2011, is now available.

Summary of Report Highlights

The *Biennial Progress Report* describes new initiatives and progress by NICEATM and ICCVAM during the period from January 2010 through December 2011. During the past two years, NICEATM, ICCVAM, and ICCVAM member agencies contributed to the national and international endorsement and adoption of 14 new and updated alternative safety testing methods. Since ICCVAM was established, NICEATM, ICCVAM, and the ICCVAM member agencies have contributed to the regulatory acceptance of over 50 alternative methods that can be used to protect and improve the health of people, animals, and the environment while reducing, refining, and replacing animal use.

Selected highlights of NICEATM and ICCVAM activities described in the *Biennial Progress Report* include:

- On behalf of NICEATM and ICCVAM, NIEHS signed an amendment to an international cooperation agreement to add the Republic of Korea and its Korean Center for the Validation of Alternative Methods (KoCVAM) to the International Cooperation on

Alternative Test Methods (ICATM). ICATM was established in 2009 by the United States, the European Union, Japan, and Canada to expedite the worldwide validation and regulatory acceptance of improved alternative test methods.

- The Organisation for Economic Co-operation and Development (OECD) adopted an international guidance document prepared by NICEATM and ICCVAM that describes how to use two cytotoxicity assays to reduce animal use for testing required to determine the poisoning potential of chemicals. NICEATM led the international validation studies for the two cytotoxicity assays, which can reduce animal use by up to 50% for each test.

- Federal agencies and the OECD adopted several new versions and applications of the murine local lymph node assay (LLNA); an alternative method recommended by ICCVAM to assess whether substances may cause allergic contact dermatitis. The test methods reduce animal use for each test by 20–40% and support expanded use of the LLNA for nearly all testing situations. Two new "green" versions of the LLNA were adopted that do not require radioactive reagents and will allow expanded use of the LLNA in laboratories worldwide.

- Federal agencies adopted ICCVAM recommended alternative test methods and procedures that will further reduce, refine, and replace animal use for eye safety testing. These include the routine use of medications to avoid most if not all pain and distress when it is necessary to use animals for required safety testing, and the first *in vitro* test method that can be used in a "bottom-up" approach to identify substances that are not considered eye hazards.

- NICEATM, ICCVAM, and their ICATM partners convened the first international workshop on alternative methods for human and veterinary vaccine potency and safety testing. The workshop reviewed the state of the science of alternative methods, and recommended priority research needed to develop improved and more efficient test methods that can also reduce, refine, and replace animal use. A focused workshop on human and veterinary rabies vaccine test methods was held in 2011 and additional focused workshops are planned for 2012 and 2013.

- ICCVAM completed international evaluation of an *in vitro* test method proposed as a screening test to identify substances with potential endocrine activity. The test method uses engineered human cells to identify substances that induce or inhibit

activation of the human estrogen receptor. Use of this test method may reduce the number of animals necessary for endocrine disruptor screening.

- NICEATM and ICCVAM convened two Best Practices for Regulatory Safety Testing Workshops to promote the use of improved and more efficient test methods that can also reduce, refine, and replace animal use. Participants learned how to select and use approved alternative methods to assess the safety or potential hazards of chemicals and products.

Background Information on ICCVAM and NICEATM

ICCVAM is an interagency committee composed of representatives from 15 Federal regulatory and research agencies that require, use, generate, or disseminate toxicological and safety testing information. ICCVAM conducts technical evaluations of new, revised, and alternative safety testing methods with regulatory applicability and promotes the scientific validation and regulatory acceptance of toxicological and safety testing methods that more accurately assess the safety and hazards of chemicals and products and that reduce, refine (enhance animal well-being and lessen or eliminate pain and distress), or replace animal use. The ICCVAM Authorization Act of 2000 (42 U.S.C. 285l-3) established ICCVAM as a permanent interagency committee of the NIEHS under NICEATM. NICEATM administers ICCVAM, provides scientific and operational support for ICCVAM-related activities, and conducts independent validation studies to assess the usefulness and limitations of new, revised, and alternative test methods and strategies. NICEATM and ICCVAM welcome the public nomination and submission of new, revised, and alternative test methods and strategies applicable to the needs of U.S. Federal agencies. Additional information about NICEATM and ICCVAM can be found on the NICEATM-ICCVAM Web site (<http://iccvam.niehs.nih.gov>).

Dated: June 4, 2012.

John R. Bucher,

Associate Director, National Toxicology Program.

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