# CONSUMER PRODUCT SAFETY COMMISSION

Notification of Request for Extension of Approval of Information Collection Requirements —Recordkeeping Requirements Under the Safety Regulations for Non-Full-Size Cribs

**AGENCY:** Consumer Product Safety Commission.

**ACTION:** Notice.

SUMMARY: In the May 18, 2004, Federal Register (69 FR 28124), the Consumer Product Safety Commission published a notice in accordance with provisions of the Paperwork Reduction Act of 1995 (44 U.S.C. chapter 35) to announce the agency's intention to seek an extension of approval of information collection requirements in the safety regulations for non-full-size cribs. 16 CFR 1500.18(a)(14) and part 1509. No comments were received on that notice. The Commission now announces that it has submitted to the Office of Management and Budget a request for extension of approval of that collection of information.

These regulations were issued to reduce hazards of strangulation, suffocation, pinching, bruising, laceration, and other injuries associated with non-full-size cribs. The regulations prescribe performance, design, and labeling requirements for non-full-size cribs. They also require manufacturers and importers of those products to maintain sales records for a period of three years after the manufacture or importation of non-full-size cribs. If any non-full-size cribs subject to provisions of 16 CFR 1500.18(a)(14) and part 1509 fail to comply in a manner severe enough to warrant a recall, the required records can be used by the manufacturer or importer and by the Commission to identify those persons and firms who should be notified of the recall.

## Additional Information About the Request for Extension of Approval of Information Collection Requirements

Agency address: Consumer Product Safety Commission, Washington, DC 20207.

Title of information collection: Recordkeeping Requirements Under the Safety Regulations for Non-Full-Size Baby Cribs, 16 CFR 1509.12.

*Type of request:* Extension of approval.

Frequency of collection: Varies, depending upon volume of products manufactured, imported, or sold.

General description of respondents: Manufacturers and importers of nonfull-size cribs. Estimated Number of respondents: 16. Estimated average number of responses per respondent: 1 per year.

Estimated number of responses for all respondents: 16 per year.

Estimated number of hours per response: 5.

*Éstimated number of hours for all respondents:* 80 per year.

Estimated cost of collection for all respondents: \$1,958.

Comments: Comments on this request for extension of approval of information collection requirements should be submitted by September 3, 2004, to (1) Office of Information and Regulatory Affairs, Attn: OMB Desk Officer for CPSC, Office of Management and Budget, Washington, DC 20503; telephone: (202) 395–7340, and (2) the Office of the Secretary, Consumer Product Safety Commission, Washington, DC 20207. Comments may also be sent to the Office of the Secretary by facsimile at (301) 504–0127 or by e-mail at cpsc-os@cpsc.gov.

Copies of this request for an extension of an information collection requirement are available from Linda L. Glatz, Office of Planning and Evaluation, Consumer Product Safety Commission, Washington, DC 20207; telephone: (301) 504–7671; or by e-mail to *lglatz@cpsc.gov*.

Dated: July 30, 2004.

## Todd A. Stevenson,

Secretary, U.S. Consumer Product Safety Commission.

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#### **DEPARTMENT OF DEFENSE**

### **Department of the Army**

Chemical and Biological Defense Program Final Programmatic Environmental Impact Statement (CBDP FPEIS)

**AGENCY:** Department of the Army, DoD. **ACTION:** Notice of availability.

SUMMARY: The Army has prepared an FPEIS covering the execution of an integrated CBDP designed to protect the members of the Armed Forces from the evolving chemical and biological (CB) threats they may encounter on the battlefield. The FPEIS includes an evaluation of how the various environmental compliance programs in the military services, the Program Executive Office for Chemical and Biological Defense, and the Defense Advanced Research Projects Agency would be able to mitigate environmental impacts.

**DATES:** The waiting period for the FPEIS will end 30 days after publication of the Notice of Availability in the **Federal Register** by the U.S. Environmental Protection Agency

ADDRESSES: Written comments or requests for copies of the FPEIS may be made to: Ms. JoLane Souris, Command Environmental Coordinator, U.S. Army Medical Research and Materiel Command, Office of Surety, Safety, and Environment, 504 Scott Street, Fort Detrick, MD 21702–5012 or visit the CBDP PEIS Web site at http://chembioeis.detrick.army.mil.

**FOR FURTHER INFORMATION CONTACT:** Ms. JoLane Souris at phone at (301) 619–2004, or by fax at (301) 619–6627.

2004, or by fax at (301) 619-6627. SUPPLEMENTARY INFORMATION: Prior to 2003, the mission of the DoD CBDP was to provide CB defense capabilities to allow the military forces of the United States to survive and successfully complete their operational missions in battle space environments contaminated with CB warfare agents. Now this mission has expanded to cover military capability to operate in the face of threats in homeland security missions, as well as war fighter missions. If our military forces are not fully and adequately prepared to meet these threats, the consequences could be devastating. The CBDP to support this mission comprises research, development, and acquisition activities. Each of the Military Services, the Joint Program Executive Office for Chemical and Biological Defense, and the Defense Advanced Research Projects Agency conduct CBDP activities. Some of these CBDP activities necessarily involve the use of hazardous chemicals or infectious disease agents for research, development, and production purposes. The controls on and the potential environmental consequences of such use for both the proposed action and the alternative were primary focuses of the CBDP FPEIS.

The activities take place at numerous military installations and contractor facilities throughout the United States. Details concerning the CBDP are contained in the Chemical and Biological Defense Program, Annual Report to Congress, April 2003 at http://www.acg.osd.mil/cp/reports.html. The proposed action consists of the execution of an integrated CBDP designed to protect the members of the Armed Forces from the evolving CB threats they may encounter on the battlefield. The No Action Alternative, continuation of current CBDP operations as described in and covered by existing environmental analyses, also was evaluated. No other alternatives