handwashing products. The new method is easy to use and inherently safe to both the user and the environment and has been submitted for patent protection.

CDC/NIOSH is soliciting for a Cooperative Research and Development Agreement (CRADA) partner to refine development of this new skin cleanser and to license and commercialize the final product. Preferred partners will have the ability to conduct testing to verify the safety of regular repeated use through a battery of clinical and instrumental test procedures aimed at determining skin compatibility. Testing trials to assess user acceptance and laboratory evaluations to determine product stability are also requested. Preferred partners should also be able to propose recommendations to the basic formulation for enhancements to user acceptance and final packaging of the end product. Preferred partners will have a strong market share and a demonstrated business network capable of effective dissemination of the final product.

Patent applications will be filed on new intellectual property resulting from the CRADA. The CRADA partner will have an option to exclusively license any rights NIOSH/CDC may have in the new technology.

Inventors: Esswein, Eric et al. U.S. Patent Application SN: Not yet filed. (CDC Ref. #: I–028–03).

Dated: August 23, 2004.

James D. Seligman,

Associate Director for Program Service, Centers for Disease Control and Prevention. [FR Doc. 04–19702 Filed 8–27–04; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Fees for Sanitation Inspections of Cruise Ships

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

ACTION: Notice.

SUMMARY: This notice announces fees for vessel sanitation inspections for fiscal year 2005 (October 1, 2004, through September 30, 2005).

EFFECTIVE DATE: October 1, 2004. **FOR FURTHER INFORMATION CONTACT:**

David L. Forney, Chief, Vessel Sanitation Program, National Center for Environmental Health, Centers for Disease Control and Prevention (CDC), 4770 Buford Highway, NE., Mailstop F–23, Atlanta, Georgia 30341–3724, telephone (770) 488–7333, e-mail: Dforney@cdc.gov.

SUPPLEMENTARY INFORMATION:

Purpose and Background

The fee schedule for sanitation inspections of passenger cruise ships inspected under the Vessel Sanitation Program (VSP) was first published in the **Federal Register** on November 24, 1987 (52 FR 45019), and CDC began collecting fees on March 1, 1988. Since then, CDC has published the fee schedule annually. This notice announces fees effective October 1, 2004.

The formula used to determine the fees is as follows: Average cost per

inspection = Total cost of VSP divided by the Weighted number of annual inspections.

The average cost per inspection is multiplied by a size/cost factor to determine the fee for vessels in each size category. The size/cost factor was established in the proposed fee schedule published in the **Federal Register** on July 17, 1987 (52 FR 27060), and revised in a schedule published in the **Federal Register** on November 28, 1989 (54 FR 48942). The revised size/cost factor is presented in Appendix A.

Fee

The fee schedule (Appendix A) will be effective October 1, 2004, through September 30, 2005. The fee schedule, which became effective October 1, 2001, will remain the same in Fiscal Year 2005. If travel expenses continue to increase, the fees may need adjustment before September 30, 2005, because travel constitutes a sizable portion of VSP's costs. If an adjustment is necessary, a notice will be published in the **Federal Register** 30 days before the effective date.

Applicability

The fees will apply to all passenger cruise vessels for which inspections are conducted as part of CDC's VSP.

Dated: August 23, 2004.

James D. Seligman,

Associate Director for Program Services, Centers for Disease Control and Prevention (CDC).

Appendix A

SIZE/COST FACTOR

Vessel size	GRT 1	Average cost (\$U.S.) per GRT
Extra Small	<3,001	0.25
Small	3,001–15,000	0.50
Medium	15,001–30,000	1.00
Large	30,001–60,000	1.50
Extra Large	>60,000	2.00

¹ Gross register tonnage in cubic feet, as shown in Lloyd's Register of Shipping.

FEE SCHEDULE OCTOBER 1, 2004—SEPTEMBER 30, 2005

Vessel size	GRT ¹	Fee
SmallMedium	<3,001 3,001–15,000 15,001–30,000 30,001–60,000	1,150 2,300 4,600 6,900

¹ Gross register tonnage in cubic feet, as shown in Lloyd's Register of Shipping.

Inspections and reinspections involve the same procedure, require the same amount of time, and are therefore charged at the same rate.

[FR Doc. 04–19703 Filed 8–27–04; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 2004D-0343]

Draft Guidance for Industry and Food and Drug Administration Staff; Hospital Bed System Dimensional Guidance to Reduce Entrapment; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug
Administration (FDA) is announcing the availability of the draft guidance entitled "Hospital Bed System
Dimensional Guidance to Reduce
Entrapment." This draft guidance provides recommendations intended to reduce life-threatening entrapments associated with hospital bed systems. It characterizes the body parts at risk for entrapment, identifies the locations of hospital bed openings that are potential entrapment areas, and recommends dimensional criteria for bed systems.

DATES: Submit written or electronic comments on this draft guidance by November 29, 2004.

ADDRESSES: Submit written requests for single copies on a 3.5" diskette of the draft guidance document entitled "Hospital Bed System Dimensional Guidance to Reduce Entrapment" to the Division of Small Manufacturers, International, and Consumer Assistance (HFZ-220), Center for Devices and Radiological Health, Food and Drug Administration, 1350 Piccard Dr., Rockville, MD 20850. Send one selfaddressed adhesive label to assist that office in processing your request, or fax your request to 301-443-8818. See the **SUPPLEMENTARY INFORMATION** section for information on electronic access to the guidance.

Submit written comments concerning this draft guidance to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.fda.gov/dockets/ecomments. Identify comments with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Jay A. Rachlin, Center for Devices and Radiological Health (HFZ–230), Food and Drug Administration, 1350 Piccard Dr., Rockville, MD 20850, 301–594–3173.

SUPPLEMENTARY INFORMATION:

I. Background

This draft guidance identifies special issues associated with hospital bed systems and provides recommendations intended to reduce life-threatening entrapments associated with these devices. Manufacturers may use this guidance to assess current hospital bed systems and to assist in the design of new beds.

II. Significance of Guidance

This draft guidance is being issued consistent with FDA's good guidance practices regulation (21 CFR 10.115). The draft guidance, when finalized, will represent the agency's current thinking on appropriate dimensional limits for gaps in hospital bed systems to prevent entrapment. It does not create or confer any rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the requirements of the applicable statute and regulations.

III. Electronic Access

To receive the "Hospital Bed System Dimensional Guidance to Reduce Entrapment" you may either send a fax request to 301–443–8818 to receive a hard copy of the document, or send an e-mail request to

GWA@CDRH.FDA.GOV to receive a hard copy or an electronic copy. Please use the document number 1537 to identify the guidance you are

requesting.

Persons interested in obtaining a copy of the draft guidance may also do so by using the Internet, CDRH maintains an entry on the Internet for easy access to information including text, graphics, and files that may be downloaded to a personal computer with Internet access. Updated on a regular basis, the CDRH home page includes device safety alerts, Federal Register reprints, information on premarket submissions (including lists of approved applications and manufacturers' addresses), small manufacturer's assistance, information on video conferencing and electronic submissions, Mammography Matters, and other device-oriented information. The CDRH Web site may be accessed at http://www.fda.gov/cdrh. A search capability for all CDRH guidance documents is available at http:// www.fda.gov/cdrh/guidance.html.

Guidance documents are also available on the Division of Dockets Management Internet site at http://www.fda.gov/ohrms/dockets.

IV. Comments

Interested persons may submit to the Division of Dockets Management (see ADDRESSES), written or electronic comments regarding this document. Submit a single copy of electronic comments or two paper copies of any mailed comments, except that individuals may submit one copy. Received comments are to be identified with the docket number found in brackets in the heading of this document. Comments received may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

Dated: August 11, 2004.

Linda S. Kahan,

Deputy Director, Center for Devices and Radiological Health.

[FR Doc. 04–19656 Filed 8–27–04; 8:45 am] **BILLING CODE 4160–01–S**

DEPARTMENT OF HOMELAND SECURITY

Bureau of Citizenship and Immigration Services

Agency Information Collection Activities: Revision of Existing Collection; Comment Request

ACTION: Request OMB Emergency Approval: Immigrant Petition for Alien Workers, 1615–0015.

The Department of Homeland Security (DHS) and the Bureau of Citizenship and Immigration Services (CIS) has submitted an emergency information collection request (ICR) utilizing emergency review procedures, to the Office of Management and Budget (OMB) for review and clearance in accordance with section 1320.13(a)(1)(ii) and (a)(2)(iii) of the Paperwork Reduction Act of 1995. The DHS has determined that it cannot reasonably comply with the normal clearance procedures under this part because normal clearance procedures are reasonably likely to prevent or disrupt the collection of information. Therefore, immediate OMB approval has been requested. If granted, the emergency approval is only valid for 180 days. ALL comments and/or questions pertaining to this pending request for emergency approval MUST be directed to OMB, Office of Information and Regulatory Affairs, Attention: Desk Officer, Department of