

Marianne Augustine at (202) 690-7102 by 5 p.m. E.D.T., September 19, 2003. Space is limited for all sessions. Written comments from the public will be accepted; opportunities to present oral comments may be provided at future meetings. Please call Marianne Augustine by 5 p.m. E.D.T., September 12, 2003, should you require a sign language interpreter. Documents pertaining to Committee deliberations will be available for public inspection and copying in Room 738-G, 200 Independence Avenue, SW., Washington, DC 20201 on the day before the meeting and following the meeting. Please call (202) 690-7102 to schedule an appointment to view the documents.

Written Comment: By this notice, the Committee is soliciting written comments, views, information and data pertinent to review of the Dietary Guidelines for Americans. Written comments are welcome throughout the Committee's deliberations. To be considered for the first meeting, they must be received by 5 p.m. E.D.T. on September 16, 2003. Comments should be sent to dietaryguidelines@osophs.dhhs.gov or to Kathryn McMurry, HHS Office of Disease Prevention and Health Promotion, Room 738-G, 200 Independence Avenue, SW., Washington, DC 20201.

Dated: August 26, 2003.

Carter Blakey,

Acting Deputy Assistant Secretary for Health, U.S. Department of Health and Human Services.

Dated: August 27, 2003.

Eric J. Hentges,

Executive Director, Center for Nutrition Policy and Promotion, U.S. Department of Agriculture.

Dated: August 28, 2003.

Edward Knipling,

Acting Administrator, Agricultural Research Service, U.S. Department of Agriculture.

[FR Doc. 03-22480 Filed 9-3-03; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Notice of Meetings

In accordance with section 10(d) of the Federal Advisory Committee Act as amended (5 U.S.C., Appendix 2), the Agency for Healthcare Research and Quality (AHRQ) announces meetings of scientific peer review groups. The subcommittees listed below are part of

the Agency's Health Services Research Initial Review Group Committee.

The subcommittee meetings will be closed to the public in accordance with the Federal Advisory Committee Act, section 10(d) of 5 U.S.C., Appendix 2 and 5 U.S.C. 552b(c)(6). Grant applications are to be reviewed and discussed at these meetings. These discussions are likely to involve information concerning individuals associated with the applications, including assessments of their personal qualifications to conduct their proposed projects. This information is exempt from mandatory disclosure under the above-cited statutes.

1. *Name of Subcommittee:* Health Care Research Training.

Date: September 25-26, 2003 (Open from 8 a.m. to 8:15 a.m. on September 25 and closed for remainder of the meeting).

Place: AHRQ Conference Center, John M. Eisenberg Bldg, Rockville, Maryland 20850.

2. *Name of Subcommittee:* Health Research Dissemination and Implementation.

Date: October 23-24, 2003 (Open from 8 a.m. to 8:15 a.m. on October 23 and closed for remainder of the meeting).

Place: AHRQ Conference Center, John M. Eisenberg Bldg, Rockville, Maryland 20850.

3. *Name of Subcommittee:* Health Care Quality and Effectiveness Research.

Date: October 23-24, 2003 (Open from 8 a.m. to 8:15 a.m. on October 23 and closed for remainder of the meeting).

Place: AHRQ Conference Center, John M. Eisenberg Bldg, Rockville, Maryland 20850.

4. *Name of Subcommittee:* Health Systems Research.

Date: October 27-28, 2003 (Open from 8 a.m. to 8:15 a.m. on October 27 and closed for remainder of the meeting).

Place: AHRQ Conference Center, John M. Eisenberg Bldg, Rockville, Maryland 20850.

5. *Name of Subcommittee:* Health Care Technology and Decision Sciences.

Date: October 30-31, 2003 (Open from 8 a.m. to 8:15 a.m. on October 30 and closed for remainder of the meeting).

Place: AHRQ Conference Center, John M. Eisenberg Bldg, Rockville, Maryland 20850.

Contact Person: Anyone wishing to obtain a roster of members, agenda or minutes of the nonconfidential portions of the meetings should contact Mrs. Bonnie Campbell, Committee Management Officer, Office of Extramural Research Review, Education and Priority Populations, AHRQ, 540 Gaither Road, Rockville, Maryland 20850, Telephone (301) 427-1554.

Agenda items for these meetings are subject to change as priorities dictate.

Dated: August 26, 2003.

Carolyn M. Clancy,

Director.

[FR Doc. 03-22476 Filed 9-3-03; 8:45 am]

BILLING CODE 4160-90-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 2003N-0376]

Medical Devices: Mammography Quality Standards Act of 1992 and Subsequent Mammography Quality Standards Reauthorization Act and Amendments; Inspection Fees

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the new fees the agency will assess for inspections of mammography facilities starting October 1, 2003. The Mammography Quality Standards Act of 1992 (the MQSA) requires FDA to assess and collect fees from mammography facilities to cover the costs of annual inspections required by the MQSA. Because these costs have increased since the last increase on February 13, 1998, FDA is raising the fees accordingly. This document explains which facilities are subject to payment of inspection fees, provides information on the costs included in developing inspection fees, and provides information on the inspection billing and collection processes. This is only the second increase in inspection fees under the MQSA since the initial fee was established in 1995.

DATES: Effective October 1, 2003, for all inspections conducted under section 354(g) of the Public Health Service Act (PHS Act) (42 U.S.C. 263b(g)). Submit written comments by October 1, 2003.

ADDRESSES: Submit written comments to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20857. 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: John L. McCrohan, Center for Devices and Radiological Health (HFZ-240), Food and Drug Administration, 1350 Piccard Dr., Rockville, MD 20850, 301-594-3332, FAX: 301-594-3306.

SUPPLEMENTARY INFORMATION:

I. Background

The MQSA requires all mammography facilities, other than facilities of the Department of Veterans Affairs, to be accredited by an approved accreditation body and certified by the

Secretary of Health and Human Services, as meeting quality standards (section 354(b) and (d)(iv) of the PHS Act). The MQSA requires FDA to establish and operate the following: (1) A Federal certification and inspection program for mammography facilities, (2) regulations and standards for accreditation bodies, and (3) standards for equipment, personnel, quality assurance, and recordkeeping and reporting by mammography facilities (section 354(c), (e), (f), and (g) of the PHS Act). The MQSA requires annual facility inspections to determine compliance with the quality standards (section 354(g) of the PHS Act). Section 354(r) of the PHS Act requires FDA to assess and collect fees for inspections of mammography facilities, other than governmental entities as determined by FDA, to cover the costs of inspections.

An updated resource review has demonstrated that the recoverable costs of the MQSA inspection program have increased since 1998. In addition, the annual amount of fees collected under the current fee schedule has been well below the level authorized by Congress. Accordingly, the fees have been recalculated so that the aggregate amount of fees collected will equal the aggregate costs of the inspections conducted, as mandated by the MQSA.

Therefore, FDA is providing notice of the increased fees to be assessed starting on October 1, 2003, and additional information relating to those fees. Although the MQSA does not require FDA to solicit comments on fee assessment and collection, FDA is inviting comments from interested persons in order to have the benefit of additional views and information, as the agency continues to evaluate its fee assessment procedures.

II. Inspections Under the Mammography Quality Standards Act of 1992

Section 354(g)(1) of the PHS Act requires FDA, States as Certifier (SAC) States, or a State or local agency acting on behalf of the FDA, to conduct an annual inspection of each mammography facility. The purpose of the annual inspection is to determine facility compliance with quality standards established under the MQSA. Inspections will be conducted by inspectors who have met Federal training requirements and who are qualified by FDA.

Under ordinary circumstances, inspections will be conducted during the regular business hours of the facility

or at a mutually agreed time. FDA normally will provide 5 working days advance notice of each annual inspection. If a significant deficiency is identified during an inspection, FDA will provide information on necessary corrective action and, in appropriate cases, will schedule a followup inspection after the facility has had a reasonable time to correct the deficiency. FDA normally will provide 5 working days advance notice of each followup inspection. FDA may make unannounced inspections or may provide shorter notice if prompt action is necessary to protect the public health (see section 354(g)(4) of the PHS Act).

III. Costs Included in the Fees to Be Assessed Beginning on October 1, 2003

Section 354(r) of the PHS Act requires FDA to assess and collect fees from persons who own or lease mammography facilities, or their agents, to cover the costs of inspections conducted by FDA, SAC States, or a State or local agency acting on behalf of FDA. Section 354(r) limits FDA's discretion in setting inspection fees in three ways: (1) Fees must be set so that, for a given fiscal year (FY), the aggregate amount of fees collected will equal the aggregate costs of inspections conducted; (2) a facility's liability for fees must be reasonably based on the proportion of the inspection costs that relate to the facility; and (3) governmental entities, as determined by FDA, are exempt from payment of fees.

FDA has determined that the following categories of costs are recoverable under section 354(r) of the PHS Act and has included them in the fees to be assessed beginning on October 1, 2003. These categories represent the same costs that have been assessed in fees since the beginning of the inspection program. Facilities are not being assessed for any new costs associated with inspections.

Cost categories are as follows: (1) Personnel costs of annual and followup inspections of mammography facilities, including administration and support; (2) purchase of equipment, calibration of instruments used in the inspections, and modification and maintenance of training facilities and laboratories to support the MQSA operations; (3) design, programming, and maintenance of data systems necessary to schedule and track inspections and to collect data during inspections; (4) training and qualification of inspectors (both FDA and State inspectors); (5) costs of billing facilities for fees due for annual and

followup inspections and collecting facility payments; (6) tracking, coordination, and direction of inspections; and (7) overhead and support attributable to facility inspections.

Because most equipment used for inspections is durable and can be used for a period of years, it is not appropriate to recover the full costs of such expenditures in the year of purchase. To do so would result in the MQSA inspection fee varying widely from one year to the next. Instead, FDA recovers these costs over the useful life of the asset.

The recoverable portions of all fixed costs of the inspection program and appropriate variable costs are recovered in the annual inspection fee. This fee will vary depending on how many mammography units are used by a facility. All mammography facilities, except governmental entities, are subject to an inspection fee.

If the annual inspection of a facility identifies a deficiency that necessitates a followup inspection that facility will be assessed an additional fee to recover the costs of that additional inspection (unless it is a governmental entity). Facilities that do not require a followup inspection are not subject to this fee.

IV. Inspection Fees to be Assessed Beginning on October 1, 2003

FDA reviewed the past methodologies for calculating the inspection fee, which accounted for differences in facility size. The same method was adopted for calculating the fees FDA will assess beginning on October 1, 2003. A facility's inspection fee will be based on the number of mammography units used by the facility.

The total recoverable aggregate cost of the MQSA inspection program is estimated to be \$14.1 million in FY 2004. This is below the \$16.4 million authorized by Congress for collections in FY 2004. To recover the costs of the inspection program, the facility portion of the fee is \$1,545 and each unit portion is \$204. The cost of each additional unit must be added to the facility portion of the fee to determine the total inspection fee. This new fee of \$1,749 for a facility with one unit compares to the current fee of \$1,549 for a facility with one unit.

FDA will assess the following fees, beginning on October 1, 2003, for facility inspections, shown in table 1 of this document:

TABLE 1.—ANNUAL INSPECTION FEE BY NUMBER OF UNITS

Number of Units	Fee
1	\$1,749
2	\$1,953
3	\$2,157
4	\$2,361
5	\$2,565
6	\$2,769
7	\$2,973
Followup Inspection Fee	\$991

FDA will continue to charge separately for annual and followup inspections. FDA believes it is more appropriate and equitable for the costs of followup inspections to be borne entirely by the facilities that require such inspections. FDA has again chosen to adopt a flat fee for followup inspections over an hourly rate that would vary the fee by the length of the inspection. This approach eliminates concerns about variations among inspectors and differential treatment of facilities. The fee schedule is subject to change each year to ensure that the aggregate amount of fees collected during any year equals the aggregate amount of costs for that year's facility inspections. FDA will monitor the adequacy of the fee on an annual basis to account for any major programmatic and budget changes.

FDA continues to use a uniform, national fee structure. The methodology adopted by FDA to determine inspection fees does not pass on the costs of inspecting governmental entities to other facilities. The entire cost of inspecting governmental entities has been and will continue to be borne by appropriated funds.

V. Facilities Subject to Payment of Inspection Fees

Under the MQSA, all certified mammography facilities except governmental entities, as determined by FDA, are subject to payment of inspection fees (see section 354(r) of the PHS Act). FDA will continue to use the definition that was previously developed and applied to determine whether a facility qualifies as a governmental entity for the purpose of determining whether a facility is exempt from payment of inspection fees under section 354(r) of the PHS Act. A facility may qualify as a governmental entity in

two ways. First, a facility may qualify if any Federal department, State, district, territory, possession, Federally-recognized Indian tribe, city, county, town, village, municipal corporation, or similar political organization does the following: (1) Operates the facility; (2) pays the entire salary of all onsite personnel for the facility; (3) owns, rents, or leases all of the facility's mammography equipment; and (4) has the ultimate authority to make day-to-day decisions concerning the management and operation of the facility.

Second, a facility may qualify as a governmental entity if the facility provides services under the Breast and Cervical Cancer Mortality Prevention Act of 1990 (www.cdc.gov/cancer/nbccedp) (FDA has verified the Web site address, but FDA is not responsible for any subsequent changes to the Web site after this document publishes in the **Federal Register**) and at least 50 percent of the mammography screening examinations provided during the preceding 12 months were funded under that statute. Facilities providing mammography services using grants under other statutes will not qualify as a government entity. FDA does not recognize, as a governmental entity, a facility providing Medicare/Medicaid services unless that facility qualifies as a governmental entity as described in the previous paragraph.

VI. Billing and Collection Procedures

Within 30 days following inspection, FDA mails a bill and a "Governmental Entity Declaration" form (Form 3422) to the inspected facility. Facilities who believe they meet the governmental entity criteria complete the form and return it in lieu of the inspection fee payment. The bill sets forth the type of inspection conducted (annual or follow-

up), the fee to be paid, and the date payment is due (30 days after billing date). Inspection fees are billed to and collected from the party that operates the facility. If the facility is owned or controlled by an entity other than the operator, it is up to the parties to establish, through contract or otherwise, how the costs of facility inspections will be allocated.

If full payment is not received by the due date, a second bill is sent. At that time, interest begins to accrue at the prevailing rate set by the Department of the Treasury, a 6 percent late payment penalty is assessed in accordance with 45 CFR 30.13, and a \$20 administrative fee is assessed for each 30-day period that a balance remains due. If payment is not received within 30 days of a third and final bill, FDA may initiate action to collect unpaid balances (with interest and penalties), including the use of collection agencies, the reporting of delinquencies to commercial credit reporting agencies, and forwarding delinquent accounts to the Department of the Treasury, Treasury Offset Program. Any questions or concerns about the billing and collection procedures may be addressed to Billing Inquiries c/o Mammography Quality Assurance Program, P.O. Box 6057, Columbia, MD 21045, 1-800-838-7715.

VII. Request for Comments

Although the MQSA does not require FDA to solicit comments on fee exemption, assessment and collection, FDA is inviting comments from interested persons in order to have the benefit of additional views.

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 paper copy. Comments are to be identified with the docket number found in brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

Dated: August 26, 2003.

Jeffrey Shuren,

Assistant Commissioner for Policy.

[FR Doc. 03-22477 Filed 9-3-03; 8:45 am]

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DEPARTMENT OF HOMELAND SECURITY

Coast Guard

DEPARTMENT OF TRANSPORTATION

Maritime Administration

[USCG-2003-14294]

El Paso Energy Bridge Gulf of Mexico, LLC Deepwater Port License Application

AGENCY: Coast Guard, DHS, and Maritime Administration, DOT.

ACTION: Notice of public hearing; request for comments.

SUMMARY: The U.S. Coast Guard (USCG) and the U.S. Maritime Administration (MARAD) will hold a public hearing to receive information relevant to the issuance or denial of the requisite federal license for the proposed El Paso Energy Bridge Gulf of Mexico, LLC (Energy Bridge GOM) Deepwater Port project. The proposed Energy Bridge GOM Deepwater Port would be located in West Cameron Area, South Addition, Block 603 (WC603) in the Central Area of the Gulf of Mexico, approximately 106 miles due south of the Louisiana coastline. We encourage interested individuals and organizations to attend the public hearing and submit comments. We also seek comments from anyone unable to attend the public hearing. In conjunction with the public hearing, the USCG and MARAD will also hold an informational open house regarding the proposed Energy Bridge GOM Deepwater Port project.

DATES: The public hearing will be held on Friday, October 3, 2003, 3 p.m. to 6 p.m., in New Orleans, Louisiana. The informational open house will be held on Friday, October 3, 2003, 1 p.m. to 3 p.m., in New Orleans, LA. The public hearing may be adjourned as early as 5 p.m. if there is no significant attendance or participation during the first two hours. The public hearing will continue

beyond 6 p.m. if necessary to ensure all individuals present at that time who wish to comment have an opportunity to do so.

Comments intended for inclusion in the public docket [USCG-2003-14294] must reach the Docket Management Facility on or before November 17, 2003.

ADDRESSES: The public hearing and informational open house will be held at the following location: New Orleans Marriott, 555 Canal Street, New Orleans, Louisiana 70130, (504) 581-1000.

You may submit comments identified by Coast Guard docket number USCG-2003-14294 to the Docket Management Facility at the U.S. Department of Transportation. To avoid duplication, please use only one of the following methods:

(1) Web Site: <http://dms.dot.gov>.

(2) Mail: Docket Management Facility, U.S. Department of Transportation, 400 Seventh Street, SW., Washington, DC 20590-0001.

(3) Fax: 202-493-2251.

(4) Delivery: Room PL-401 on the Plaza level of the Nassif Building, 400 Seventh Street, SW., Washington, DC, between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays. The telephone number is 202-366-9329.

(5) Federal eRulemaking Portal: <http://www.regulations.gov>.

FOR FURTHER INFORMATION CONTACT: For further information concerning this notice, the Energy Bridge GOM Deepwater Port license application, or the public hearing or informational open house, contact Commander Mark Prescott, U.S. Coast Guard at (202) 267-0225 or mprescott@comdt.uscg.mil. For questions on viewing or submitting material to the docket, call Ms. Andrea Jenkins, Program Manager, Docket Operations, Department of Transportation, at (202) 366-0271.

SUPPLEMENTARY INFORMATION:

Public Participation and Request for Comments

Whether or not you attend the public hearing or informational open house, we encourage you to submit written comments and related materials. All comments received will be posted, without change, to <http://dms.dot.gov> and will include any personal information you have provided. We have an agreement with the Department of Transportation (DOT) to use the Docket Management Facility. Please see DOT's "Privacy Act" paragraph below.

Submitting comments: If you submit a comment, please include your name and address, identify the docket number

[USCG-2003-14294], indicate your specific concern, and give the reason for each comment. You may submit your comments and material by electronic means, mail, fax, or delivery to the Docket Management Facility at the address under **ADDRESSES**; but please submit your comments and material by only one means. If you submit them by mail or delivery, submit them in an unbound format, no larger than 8½ by 11 inches, suitable for copying and electronic filing. If you submit them by mail and would like to know that they reached the Facility, please enclose a stamped, self-addressed postcard or envelope. We will consider all comments and material received during the comment period.

Viewing comments and documents:

To view comments, as well as documents mentioned in this preamble as being available in the docket, go to <http://dms.dot.gov> at any time and conduct a simple search using the docket number USCG-2003-14294. You may also visit the Docket Management Facility in room PL-401 on the Plaza level of the Nassif Building, 400 Seventh Street, SW., Washington, DC, between 9 a.m. and 5 p.m., Monday through Friday, except Federal holidays.

Privacy Act: Anyone can search the electronic form of all comments received into any of our dockets by the name of the individual submitting the comment (or signing the comment, if submitted on behalf of an association, business, labor union, etc.). You may review the Department of Transportation's Privacy Act Statement in the **Federal Register** published on April 11, 2000 (65 FR 19477), or you may visit <http://dms.dot.gov>.

Public Hearing/Informational Open House

The Coast Guard and the Maritime Administration will hold a public hearing from 3 p.m. to 6 p.m. on Friday, October 3, 2003, at the New Orleans Marriott, 555 Canal Street, New Orleans, Louisiana. An informational open house will be held prior to the public meeting from 1 p.m. to 3 p.m. at the same location. We invite the public and representatives of interested agencies to attend and provide comments on the proposed license application. If you plan to attend the public hearing or informational open house and need special assistance, such as sign language interpretation or other reasonable accommodations, contact the U.S. Coast Guard as indicated in **FOR FURTHER INFORMATION CONTACT**.