This section of the FEDERAL REGISTER contains notices to the public of the proposed issuance of rules and regulations. The purpose of these notices is to give interested persons an opportunity to participate in the rule making prior to the adoption of the final rules.

DEPARTMENT OF COMMERCE
Office of the Secretary
15 CFR Part 4
[Docket No. 150324296–5527–02]
RIN 0605–AA38

Public Information, Freedom of Information Act and Privacy Act

AGENCY: Office of the Secretary, U.S. Department of Commerce.

ACTION: Proposed rule; reopening of public comment period.

SUMMARY: On May 8, 2015, the Department of Commerce (Department) published in the Federal Register a proposed rule to revise the Department’s regulations under the Privacy Act. In particular, the Department proposed to amend its regulations regarding applicable exemptions to the Privacy Act to reflect new Department wide systems of records notices published since the last time the regulations were updated. The Department opened a public comment period through June 8, 2015. The Department is reopening the original public comment period of 30 days for the proposed rulemaking for an additional 30 days from the date of publication of this notice. The reopening is necessary because not all interested parties may have been given appropriate notification about this proposed new system of records, as well as time to respond with comments prior to the closing date of the original public comment period of June 8, 2015.

DATES: The comment period for the proposed rule published May 8, 2015 (80 FR 26499), is reopened. To be considered, written comments must be submitted on or before July 29, 2015.

ADDRESSES: You may submit written comments, identified by Regulatory Identification Number (RIN) 0605–AA38, by any of the following methods:

• Federal eRulemaking Portal: http://www.regulations.gov. Follow the instructions for submitting comments.

• Fax: (202) 482–0827. Include RIN 0605–AA38 in the subject line.

• Mail: Dr. Michael Toland, Office of Privacy and Open Government, 1401 Constitution Ave. NW., Washington, DC 20230.

Instructions: All submissions received must include the agency name and docket number or RIN for this rulemaking. All comments received will be posted without change to regulations.gov, including any personal information received. For detailed instructions on submitting comments and additional information on the rulemaking process, see the “Public Participation” heading of the SUPPLEMENTARY INFORMATION section for the proposed rule published May 8, 2015 (80 FR 26499).


SUPPLEMENTARY INFORMATION: On May 8, 2015, the Department of Commerce published in the Federal Register a rule proposing revisions to the Department’s Privacy Act regulations under the Privacy Act. The revisions of the Privacy Act regulations in subpart B of part 4 incorporate changes to the language in the regulations in the following provisions: § 4.33 (General exemptions); and § 4.34 (Specific Exemptions). See 80 FR 26499, May 8, 2015. In the original proposal, the Department requested comment on or before June 8, 2015. The public comment period is being reopened because not all interested parties may have been given adequate notice, as well as time to respond with comments about the Department’s proposal to revise its regulations under the Privacy Act of 1974. This reopening will provide an opportunity for appropriate review and comment of the proposed rulemaking now posted in regulations.gov, RIN 0605–AA38, or in the public docket, Docket ID No. 150324296–5265–01, for 30 days from the date of today’s publication.

All previously submitted comments will be responded to as appropriate, and members of the public who have submitted comments during the prior comment period need not resubmit them at this time. Comments received after June 8, but before publication of this notice must be resubmitted in order to be considered.

If no comments are received, a rule will be published, effective as proposed in 80 FR 26499 (May 8, 2015) on the date of publication, in the Federal Register.

Dated: June 19, 2015.

Catrina D. Purvis, Department of Commerce, Chief Privacy Officer and Director for Open Government.

[FR Doc. 2015–15865 Filed 6–26–15; 8:45 am]

BILLING CODE 3510–BX–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 2
[Docket No. FDA–2015–N–1355]

Use of Ozone-Depleting Substances; Request for Comment Concerning Essential-Use Designations

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice and request for public comment.

SUMMARY: The Food and Drug Administration (FDA or Agency) is seeking public comment on whether the uses of ozone-depleting substances (ODSs), including chlorofluorocarbons (CFCs), in certain FDA-regulated products currently designated essential are no longer essential under the Clean Air Act due to the availability of alternatives that do not use CFCs or because the products are no longer being marketed. Essential-use products are exempt from FDA’s ban on the use of CFC propellants in FDA-regulated products and the Environmental Protection Agency’s (EPA’s) ban on the use of CFCs in pressurized dispensers. FDA is seeking public comment because it is responsible for determining which FDA-regulated products that release CFCs or other ODSs are essential uses under the Clean Air Act. FDA is soliciting comments to assist the Agency in striking an appropriate balance that will best protect the public health, both by ensuring the availability of an adequate number of alternatives and by curtailing the release of ODSs.

DATES: Submit written or electronic comments by August 28, 2015.
provide unique health benefits that would not be available without the use of a CFC.

EPA regulations implementing the provisions of section 610 of the Clean Air Act (42 U.S.C. 7671i) contain a general ban on the use of CFCs in pressurized dispensers, such as metered-dose inhalers (MDIs) (40 CFR 82.64(c) and 82.66(d)). These EPA regulations exempt from the general ban “medical devices” that FDA considers essential and that are listed in 21 CFR 2.125(e). Section 2.125(b) of the Clean Air Act (42 U.S.C. 7671(b)) defines “medical device” as any device (as defined in the FD&C Act), diagnostic product, drug (as defined in the FD&C Act), and drug delivery system, if such device, diagnostic product, drug, or drug delivery system uses a class I or class II ODS for which no safe and effective alternative has been developed (and, where necessary, approved by the Commissioner of Food and Drugs (the Commissioner)); and if such device, diagnostic product, drug, or drug delivery system is considered essential. Section 601(8) of the Clean Air Act (section 601 (8) of the Clean Air Act (42 U.S.C. 7671(8)) contains a “medical device” as any device (as defined in the FD&C Act), diagnostic product, drug, or drug delivery system, if such device, diagnostic product, drug, or drug delivery system uses a class I or class II ODS for which no safe and effective alternative has been developed (and, where necessary, approved by the Commissioner of Food and Drugs (the Commissioner)); and if such device, diagnostic product, drug, or drug delivery system is considered essential. Section 2.125(g) of the Clean Air Act (42 U.S.C. 7671(g)) provides a list of medical devices (as defined in section 201(h) of the Clean Air Act) covered under section 610 of the Clean Air Act must receive exemptions for essential uses under the Montreal Protocol.

Faced with the statutorily mandated phase-out of the production of CFCs, drug manufacturers have developed alternatives to MDIs and other self-pressurized drug dosage forms that do not contain ODSs. Examples of these alternative dosage forms are MDIs that use non-ODSs as propellants and dry-powder inhalers. The current and future availability of technically feasible alternatives to the use of a CFC may mean that the existing listing of an essential use in 21 CFR 2.125(e) no longer reflects current conditions. It is with this situation in mind that FDA is seeking public comment regarding the Agency’s consideration of whether certain uses of ODSs are no longer essential.

FDA is soliciting comments to assist the Agency in striking an appropriate balance that will best protect the public health by ensuring the availability of an adequate number of treatment alternatives and by curtailing the release of ODSs.

II. Listed Uses That May No Longer Be Considered Essential

There are certain uses of CFCs currently listed in 21 CFR 2.125(e) as essential that may no longer be considered as essential. Section 2.125(g) sets forth standards for determining whether the use of an ODS in a medical product is no longer essential. FDA is seeking public comment concerning whether the following uses should still be considered essential:

A. Sterile Aerosol Talc Administered Intrapleurally by Thoracoscopy for Human Use (21 CFR 2.125(e)(4)(ix))

There is currently one sterile aerosol talc product containing ODSs which is approved for administration intrapleurally by thoracoscopy for human use for the treatment of recurrent symptomatic malignant pleural effusion (MPE) in symptomatic patients. Under 21 CFR 2.125(g)(3), an essential-use designation for individual active moieties marketed as ODS products and represented by a new drug application may no longer be essential if:

- At least one non-ODS product with the same active moiety is marketed with the same route of administration, for the same indication, and with approximately the same level of convenience of use as the ODS product containing that active moiety;
- Supplies and production capacity for the non-ODS product(s) exist or will exist at levels sufficient to meet patient need;
• Adequate U.S. postmarketing-use data are available for the non-ODS product(s); and
• Patients who medically require the ODS product are adequately served by the non-ODS product(s) containing that active moiety and other available products (21 CFR 2.125(g)(3)).

Sterile aerosol talc is currently marketed for intrapleural administration in two non-ODS formulations—powder and aerosol. Sterile aerosol talc is a powder formulation of talc available for intrapleur al administration via chest tube. Sclerosol Intrapleural Aerosol (sterile talc powder) is an aerosol formulation which contains the propellant, hydrofluoroalkane (HFA) 134a and is approved for intrapleural administration. Sclerosol Intrapleural Alcohol, a form of aerosol sterile talc, is indicated for the treatment of recurrent MPE in symptomatic patients.

The route of administration, indications, and level of convenience appear to be the same for the ODS and non-ODS formulations of sterile aerosol talc. Moreover, because production of non-ODS formulations are not limited by restrictions on the use of ODSs, the Agency believes that non-ODS formulations can be produced at greater quantities and have the potential to be more widely available than prior formulations that contained ODSs. In addition, there is adequate U.S. postmarketing-use data indicating that the non-ODS products are available in sufficient quantities to serve the current patient population. For these reasons, we believe that patients may be adequately served by the non-ODS products containing sterile aerosol talc. Thus, FDA is seeking public comment concerning whether sterile aerosol talc administered intrapleurally by thoracoscopy for human use is no longer an essential use of ODSs described in 21 CFR 2.125(e).

B. Drug Products That Are No Longer Being Marketed

Under 21 CFR 2.125(g)(1), an active moiety may no longer be an essential-use (21 CFR 2.125(e)) if it is no longer marketed in an approved ODS formulation. FDA believes failure to market indicates non-essentiality because the absence of a demand sufficient for even one company to market the product is highly indicative that the use is not essential.

FDA is seeking public comment as to whether metered-dose atropine sulfate aerosol human drugs administered by oral inhalation (21 CFR 2.125(e)(4)(i)) and anesthetic drugs for topical use on accessible mucus membranes of humans where a cannula is used for application (21 CFR 2.125(e)(4)(iii)) are no longer essential uses as described at 21 CFR 2.125(e). FDA has information that these products are not currently being marketed in an approved form that releases ODSs, and, under 21 CFR 2.125(g)(1), they may no longer constitute an essential-use. Because these products are no longer being marketed, FDA does not believe that loss of essential use status would not result in any drugs being made unavailable to patients.

III. Comments

Interested persons may submit either electronic comments regarding this document to http://www.regulations.gov or written comments to the Division of Dockets Management (see ADDRESSES). It is only necessary to send one set of comments. Identify comments 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, and will be posted to the docket at http://www.regulations.gov.

Dated: June 24, 2015.
Leslie Kux,
Associate Commissioner for Policy.

[FR Doc. 2015–15902 Filed 6–26–15; 8:45 am]
BILLING CODE 4164–01–P

DEPARTMENT OF TRANSPORTATION
Federal Highway Administration

23 CFR Parts 630 and 635
[Docket No. FHWA–2015–0009; FHWA RIN 2125–AF61
Construction Manager/General Contractor Contracting

AGENCY: Federal Highway Administration, DOT.

ACTION: Notice of proposed rulemaking (NPRM); request for comments.

SUMMARY: Section 1303 of the Moving Ahead for Progress in the 21st Century Act (MAP–21) amends 23 U.S.C. 112 to require the Secretary of Transportation to promulgate regulations as necessary to implement the Construction Manager/General Contractor (CM/GC) contracting method. This NPRM initiates the formal rulemaking process to fulfill the legislative requirement and establish such regulations as are necessary for the FHWA’s approval of projects using the CM/GC method of contracting.

DATES: Comments must be received on or before August 28, 2015.

ADDRESSES: To ensure that you do not duplicate your docket submissions, please submit them by only one of the following means:

• Federal eRulemaking Portal: Go to http://www.regulations.gov and follow the online instructions for submitting comments.
• Mail: Docket Management Facility, U.S. Department of Transportation, 1200 New Jersey Ave. SE., West Building Ground Floor Room W12–140, Washington, DC 20590–0001;
• Hand Delivery: West Building Ground Floor, Room W12–140, 1200 New Jersey Ave. SE., between 9 a.m. 5 p.m., Monday through Friday, except Federal holidays. The telephone number is (202) 366–9329:

• Instructions: You must include the agency name and docket number DOT–FHWA–or the Regulatory Identification Number (RIN) for the rulemaking at the beginning of your comments. All comments received will be posted without change to http://www.regulations.gov, including any personal information provided.


U.S. Department of Transportation, 1200 New Jersey Avenue SE., Washington, DC 20590. Office hours are from 8 a.m. to 4:30 p.m., E.T., Monday through Friday, except Federal holidays.

SUPPLEMENTARY INFORMATION:

Summary

This regulatory action is undertaken to fulfill the statutory requirement in Section 1303(b) of MAP–21 requiring the Secretary to promulgate a regulation to implement the CM/GC method of contracting. The CM/GC is a contracting method that allows a contracting agency to use a single procurement to secure pre-construction and construction services. In the pre-construction services phase, a contracting agency procures the services of a construction contractor early in the design phase of a project in order to obtain the contractor’s input on constructability issues that may be affected by the project design. A CM/GC contractor does not provide any preliminary or final design services. As part of the preconstruction services phase of a CM/GC contract, the CM/GC contractor provides information for consideration in the design and environmental review processes on construction-related aspects of a project, including the potential effects of design elements on...