cause for the extension. The FAA also has determined that extension of the comment period is consistent with the public interest, and that good cause exists for taking this action.

Accordingly, the comment period for the Repair Stations, NPRM, Docket No. FAA-2006-26408, is extended until April 16, 2007.

Issued in Washington, DC, February 20, 2007.

## James J. Ballough,

Director, Flight Standards Service, Aviation Safety.

[FR Doc. E7–3331 Filed 2–26–07; 8:45 am] BILLING CODE 4910–13–P

### RAILROAD RETIREMENT BOARD

#### 20 CFR Part 230

RIN 3220-AA61

# Reduction and Nonpayment of Annuities by Reason of Work

**AGENCY:** Railroad Retirement Board. **ACTION:** Proposed rule; withdrawal.

**SUMMARY:** The above mentioned regulation was previously published as a proposed rule on August 16, 1995 (60 FR 42482). The Railroad Retirement Board has determined not to go final with that proposed rule and hereby withdraws the proposed rule to amend 20 CFR Part 230.

**ADDRESSES:** 844 North Rush Street, Chicago, Illinois 60611–2092.

# FOR FURTHER INFORMATION CONTACT:

Marguerite P. Dadabo, Assistant General Counsel, Office of General Counsel, Railroad Retirement Board, (312) 751– 4945, FAX (312) 751–7102, TDD (312) 751–4701.

Dated: February 21, 2007.

## Beatrice Ezerski,

Secretary to the Board.

[FR Doc. 07–872 Filed 2–26–07; 8:45 am]

BILLING CODE 7905-01-M

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# **Food and Drug Administration**

## 21 CFR Part 868

[Docket No. 2007N-0019]

Medical Devices; Anesthesiology Devices; Oxygen Pressure Regulators and Oxygen Conserving Devices

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Proposed rule.

**SUMMARY:** The Food and Drug Administration (FDA) is publishing a proposed rule to reclassify pressure regulators for use with medical oxygen, currently class I devices included in the generic type of device called pressure regulator, into class II, subject to special controls in the form of a guidance document. Pressure regulators for use with all other medical gases will remain in class I, subject only to general controls. FDA is also proposing to establish a separate classification regulation for oxygen conserving devices (or oxygen conservers), now included in the generic type of device called noncontinuous ventilator. Oxygen conserving devices will continue to be classified in class II, but those that incorporate a built-in oxygen pressure regulator will become subject to the special controls guidance if the rule is finalized. Elsewhere in this issue of the **Federal Register**, FDA is announcing the availability of a class II special controls draft guidance for industry and FDA staff entitled "Class II Special Controls Guidance Document: Oxygen Pressure Regulators and Oxygen Conserving Devices." The agency is proposing this action because it believes that special controls are necessary to provide a reasonable assurance of safety and effectiveness for these devices.

**DATES:** Submit comments by May 29, 2007. FDA is proposing that any final rule based on this proposed rule be effective 2 years after the date of its publication in the **Federal Register**.

ADDRESSES: You may submit comments, identified by Docket No. 2007N–0019, by any of the following methods: *Electronic Submissions*Submit electronic comments in the following ways:

- Federal eRulemaking Portal: http://www.regulations.gov. Follow the instructions for submitting comments.
- Agency Web site: http:// www.fda.gov/dockets/ecomments. Follow the instructions for submitting comments on the agency Web site. Written Submissions Submit written submissions in the following ways:
  - FAX: 301-827-6870.
- Mail/Hand delivery/Courier [For paper, disk, or CD–ROM submissions]: Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

To ensure more timely processing of comments, FDA is no longer accepting comments submitted to the agency by email. FDA encourages you to continue to submit electronic comments by using the Federal eRulemaking Portal or the agency Web site, as described in the **ADDRESSES** portion of this document under *Electronic Submissions*.

Instructions: All submissions must include the agency name and Docket No. for this rulemaking. All comments received may be posted without change to <a href="http://www.fda.gov/ohrms/dockets/default.htm">http://www.fda.gov/ohrms/dockets/default.htm</a>, including any personal information provided. For additional information on submitting comments, see section XII "What if I Have Comments to the Proposed Rule" heading in the SUPPLEMENTARY INFORMATION section of this document.

Docket: For access to the docket to read background documents or comments received, go to http://www.fda.gov/ohrms/dockets/default.htm and insert the docket number, found in brackets in the heading of this document, into the "Search" box and follow the prompts and/or go to the Division of Dockets Management, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

## FOR FURTHER INFORMATION CONTACT:

Christy Foreman, Center for Devices and Radiological Health (HFZ–340), Food and Drug Administration, 2094 Gaither Rd., Rockville, MD 20850, 240–276– 0120.

#### SUPPLEMENTARY INFORMATION:

# I. What Are the Highlights of the Proposed Rule?

The highlights of the proposed rule are as follows:

- FDA is dividing the classification of pressure regulators into two classification regulations.
- Pressure regulators for use with medical gases other than oxygen will remain in class I.
- Pressure regulators for use with medical oxygen will be identified as "oxygen pressure regulators" and will be reclassified into class II (special controls).
- FDA is establishing a separate classification regulation for oxygen conserving devices, which are now included in the generic type of device called noncontinuous ventilators.
- Both noncontinuous ventilators and oxygen conserving devices will remain in class II.
- Oxygen conservers will be classified within their own class according to whether or not the device incorporates a built-in oxygen pressure regulator.
- FDA is establishing a special controls guidance document for oxygen pressure regulators and oxygen conservers that have built-in oxygen pressure regulators entitled "Class II Special Controls Guidance Document: Oxygen Pressure Regulators and Oxygen