necessary for the safety of the spacecraft and personnel on board, subject any of the personnel on board to such restraint as the circumstances require until such time as delivery of such individual or individuals to the proper authorities is possible.

10. Amend paragraphs (a), (c) and (d) in § 1214.703 to read as follows:

§ 1214.703 Chain of command.

(a) The NASA Commander is a trained NASA astronaut who has been designated to serve as commander on a NASA mission and who shall have the authority described in § 1214.702 of this part. Under normal flight conditions (other than emergencies or when otherwise designated) the NASA Commander is responsible to the Mission Flight Director.

(c) Before each flight, the other flight crewmembers will be designated in the order in which they will assume the authority of the NASA Commander under this subpart in the event that the NASA Commander is not able to carry out his/her duties.

(d) The determinations, if any, that a crewmember in the chain of command is not able to carry out his or her command duties and is, therefore, to be relieved of command, and that another crewmember in the chain of command is to succeed to the authority of the NASA Commander, will be made by the NASA Administrator or his/her designee.

11. Revise § 1214.704 to read as follows:

§ 1214.704 Violations.

(a) All personnel on board the NASA mission are subject to the authority of the NASA Commander and shall conform to his/her orders and direction as authorized by this subpart.

(b) This regulation is a regulation within the meaning of 18 U.S.C. 799, and whoever willfully violates, attempts to violate, or conspires to violate any provision of this subpart or any order or direction issued under this subpart shall be subject to fines and imprisonment, as specified by law.

Subpart 1214.8—[Removed and Reserved]

12. Remove and reserve subpart 1214.8, consisting §§ 1214.800 through 1214.813.

Subpart 1214.17—[Removed and Reserved]

13. Remove and reserve subpart 1214.17, consisting of §§ 1214.1700 through 1214.1707.

Nanette Jennings,
Federal Register Liaison Officer.
[FR Doc. 2015–26475 Filed 10–19–15; 8:45 am]
BILLING CODE 7510–13–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration
21 CFR Part 101
[Docket No. FDA–2012–N–1210]
RIN 0910–AF22

Food Labeling: Revision of the Nutrition and Supplement Facts Labels; Reopening of the Comment Period

AGENCY: Food and Drug Administration, HHHS.

ACTION: Proposed rule; reopening of comment period.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing the reopening of the comment period for certain documents associated with the proposed rule to amend FDA’s labeling regulations for conventional foods and dietary supplements to provide updated nutrition information on the Nutrition Facts and Supplement Facts labels to assist consumers in maintaining healthy dietary practices. We also are reopening the comment period for a supplemental proposed rule to revise the Nutrition Facts and Supplement Facts labels. We are taking this action due to technical difficulties at the Federal eRulemaking Portal.

DATES: Submit either electronic or written comments on the supplemental proposed rule and related documents by October 23, 2015.

ADDRESSES: You may submit comments as follows:

Electronic Submissions

Submit electronic comments in the following way:

• Federal eRulemaking Portal: http://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to http://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on http://www.regulations.gov.

• If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see “Written/Paper Submissions” and “Instructions”).

Written/Paper Submissions

Submit written/paper submissions as follows:

Mail/Hand delivery/Courier (for written/paper submissions): Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fithers Lane, Rm. 1061, Rockville, MD 20852.

• For written/paper comments submitted to the Division of Dockets Management, FDA will post your comment, as well as any attachments, except for information submitted, marked and identified, as confidential, if submitted as detailed in “Instructions.”

Instructions: All submissions received must include the Docket No. FDA–2012–N–1210 for this rulemaking. Received comments will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at http://www.regulations.gov or at the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

• Confidential Submissions—To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states “THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION”. The Agency will review this copy, including the claimed confidential information, in its consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on http://www.regulations.gov. Submit both copies to the Division of Dockets Management. If you do not wish your
name and contact information to be made publicly available, you can provide this information on the cover sheet and not in the body of your comments and you must identify this information as “confidential.” Any information marked as “confidential” will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA’s posting of comments to public dockets, see 80 FR 56469, September 18, 2015, or access the information at: http://www.fda.gov/regulatoryinformation/dockets/default.htm.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to http://www.regulations.gov 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:

SUPPLEMENTARY INFORMATION:
I. Background

In the Federal Register of March 3, 2014 (79 FR 11879), we published a proposed rule that would amend our labeling regulations for conventional foods and dietary supplements to provide updated nutrition information. In the Federal Register of July 27, 2015 (80 FR 44302), we reopened the comment period through September 25, 2015, for the proposed rule for the sole purpose of inviting public comments on two consumer studies being added to the administrative record. The consumer studies pertained to proposed changes to the Nutrition Facts label formats. We also issued a supplemental proposed rule (80 FR 44303) with a comment period through October 13, 2015. The supplemental proposal included two additional consumer studies pertaining to the declaration of added sugars and alternative footnote statements. We proposed text for the footnotes to be used on the Nutrition Facts label, after completing our consumer research in which we tested various footnote text options for the label. We also proposed to establish a Daily Reference Value of 10 percent of total energy intake from added sugars and to require the declaration of the percent Daily Value for added sugars on the label. The supplemental proposed rule also provided additional rationale for the declaration of the amount of added sugars on the label. We explained that we were taking these actions based, in part, on the science underlying a new report released by the 2015 Dietary Guidelines Advisory Committee.

More recently, in the Federal Register of September 10, 2015 (80 FR 54446), we issued a notice clarifying: (1) The consumer studies on the added sugars declaration and the alternative footnote statements in the supplemental proposal relate to topics on which we sought comment and (2) the consumer studies on the format published in a separate notice in July 2015 were included for comment, and were placed in the docket at that time. We also stated that, in response to requests for the raw data for each of these consumer studies that are relevant to the summary memorandum for the studies, we were making the raw data available for comment. We extended the comment period for the two consumer studies pertaining to the proposed changes to the Nutrition Facts label formats (originally scheduled to close on September 25, 2015) to October 13, 2015, to coincide with the end of the comment period for the supplemental proposed rule.

However, on October 13 and 14, 2015, the Federal eRulemaking Portal, http://www.regulations.gov, experienced technical difficulties which sometimes prevented the electronic submission of comments. Therefore, we are reopening the comment period for the consumer studies and the supplemental proposal; the reopened comment period will close on October 23, 2015.

Leslie Kux,
Associate Commissioner for Policy.

BILLING CODE 4164–01–P

DEPARTMENT OF EDUCATION
34 CFR Chapter VI
[Docket ID ED–2015–OPE–0103]

Negotiated Rulemaking Committee; Negotiator Nominations and Schedule of Committee Meetings—Borrower Defenses

AGENCY: Office of Postsecondary Education, Department of Education.

ACTION: Intent to establish negotiated rulemaking committee.

SUMMARY: We announce our intention to establish a negotiated rulemaking committee to prepare proposed regulations for the Federal Student Aid programs authorized under title IV of the Higher Education Act of 1965, as amended (HEA). The committee will include representatives of organizations or groups with interests that are significantly affected by the topics proposed for negotiations. We request nominations for individual negotiators who represent key stakeholder constituencies for the issues to be negotiated to serve on the committee, and we set a schedule for committee meetings.

DATES: We must receive your nominations for negotiators to serve on the committee on or before November 19, 2015. The dates, times, and locations of the committee meetings are set out in the Schedule for Negotiations section in the SUPPLEMENTARY INFORMATION section.


FOR FURTHER INFORMATION CONTACT: For information about the content of this notice, including information about the negotiated rulemaking process or the nomination submission process, contact: Wendy Macias, U.S. Department of Education, 1990 K Street NW., Room 8013, Washington, DC 20006. Telephone: (202) 502–7526 or by email: Wendy.Macias@ed.gov.


If you use a telecommunications device for the deaf (TDD) or text telephone (TTY), call the Federal Relay Service (FRS) toll free at 1–800–877–8339.

SUPPLEMENTARY INFORMATION: On August 20, 2015, we published a notice in the Federal Register (80 FR 50588) announcing our intent to establish a negotiated rulemaking committee under section 492 of the HEA to develop proposed regulations for determining which acts or omissions of an institution of higher education (“institution”) a borrower may assert as a defense to repayment of a loan made under the William D. Ford Federal Direct Loan (Federal Direct Loan) Program (“borrower defenses”) and the consequences of such borrower defenses for borrowers, institutions, and the