must be tested (Aviation flight crew members and air traffic controllers);
(c) An occupation identified in 49 CFR 382.103 by the Federal Motor Carrier Safety Administration, in which the employee must be tested (Commercial drivers);
(d) An occupation identified in 49 CFR 219.3 by the Federal Railroad Administration, in which the employee must be tested (Railroad operating crew members);
(e) An occupation identified in 49 CFR 653.5 by the Federal Transit Administration, in which the employee must be tested (Public transportation operators);
(f) An occupation identified in 49 CFR 199.2 by the Pipeline and Hazardous Materials Safety Administration, in which the employee must be tested (Pipeline operation and maintenance crew members);
(g) An occupation identified in 49 CFR 16.201 by the United States Coast Guard, in which the employee must be tested (Crewmembers and maritime credential holders on a commercial vessel);
(b) An occupation specifically identified in Federal law as requiring an employee to be tested for controlled substances;
(i) An occupation specifically identified in the State law of that State as requiring an employee to be tested for controlled substances; and
(j) An occupation where the State has a factual basis for finding that employers hiring employees in that occupation conduct pre- or post-hire drug testing as a standard eligibility requirement for obtaining or maintaining employment in the occupation.

§ 620.4 Testing of unemployment compensation applicants for the unlawful use of a controlled substance.

(a) States may require drug testing for unemployment compensation applicants, as defined in § 620.2, for the unlawful use of one or more controlled substances, as defined in § 620.2, as a condition of eligibility for unemployment compensation, if the individual is one for whom suitable work, as defined in State law, as defined in § 620.2 of, is only available in an occupation that regularly conducts drug testing as identified under § 620.3.
(b) A State conducting drug testing as a condition of unemployment compensation eligibility, as provided in paragraph (a) of this section, may only elect to require drug testing of applicants for whom the only suitable work is available in one or more of the occupations listed under § 620.3. States are not required to apply drug testing to any applicants for whom the only suitable work is available in any or all of the occupations listed.
(c) No State is required to drug test UC applicants under this part 620.

§ 620.5 Conformity and substantial compliance.

(a) In general. A State law implementing the drug testing of applicants for unemployment compensation must conform with—and the law’s administration must substantially comply with—the requirements of this part 620 for purposes of certification under 42 U.S.C. 502(a), governing State eligibility to receive Federal grants for the administration of its UC program.
(b) Resolving Issues of Conformity and Substantial Compliance. For the purposes of resolving issues of conformity and substantial compliance with the requirements of this part 620, the provisions of 20 CFR 601.5 apply.

Molly E. Conway,
Acting Assistant Secretary for Employment and Training, Labor.

[FR Doc. 2018–23952 Filed 11–2–18; 8:45 am]
BILLING CODE 4510–FW–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 15
[Docket No. FDA–2018–N–3952]

Eliminating Youth Electronic Cigarette and Other Tobacco Product Use: The Role for Drug Therapies; Public Hearing; Request for Comments

AGENCY: Food and Drug Administration, HHHS.
ACTION: Notification of public hearing; request for comments.

SUMMARY: The Food and Drug Administration (FDA or the Agency) is announcing a public hearing to discuss its efforts to eliminate youth electronic cigarette (e-cigarette) use as well as other tobacco product use, with a focus on the potential role of drug therapies to support youth e-cigarette cessation and the issues impacting the development of such therapies.

DATES: The public hearing will be held on December 5, 2018, from 9 a.m. to 5 p.m. The public hearing may be extended or may end early depending on the level of public participation.

Persons seeking to present at the public hearing must register by Friday, November 23, 2018. Persons seeking to attend, but not present at, the public hearing must register by Monday, December 3, 2018. Section II provides attendance and registration information. Electronic or written comments will be accepted after the public hearing until Wednesday, January 2, 2019.

ADDRESSES: The public hearing will be held at the FDA White Oak Campus, 10903 New Hampshire Ave., Building 31 Conference Center, the Great Room A, Silver Spring, MD 20993–0002. Entrance for public hearing participants (non-FDA employees) is through Building 1, where routine security check procedures will be performed. For parking and security information, please refer to https://www.fda.gov/AboutFDA/WorkingatFDA/BuildingsandFacilities/WhiteOakCampusInformation/ucm241740.htm.

You may submit comments as follows. Please note that late, untimely filed comments will not be considered. Electronic comments must be submitted on or before Wednesday, January 2, 2019. The https://www.regulations.gov electronic filing system will accept comments until 11:59 p.m. Eastern Time at the end of Wednesday, January 2, 2019. Comments received by mail/hand delivery/courier (for written/paper submissions) will be considered timely if they are postmarked or the delivery service acceptance receipt is on or before that date. You may submit comments as follows:

Electronic Submissions

Submit electronic comments in the following way:

• Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://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 https://www.regulations.gov.

• If you want to submit a comment with a 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): Dockets Management Staff (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.
- For written/paper comments submitted to the Dockets Management Staff, 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–2018–N–3952 for “Eliminating Youth Electronic Cigarette and Other Tobacco Product Use: The Role for Drug Therapies; Public Hearing; Request for Comments.” Received comments will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at https://www.regulations.gov or at the Dockets Management Staff 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 https://www.regulations.gov. Submit both copies to the Dockets Management Staff. 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 docket, see 80 FR 56409, September 18, 2015, or access the information at: https://www.fda.gov/regulatoryinformation/dockets/default.htm.

Docket: For access to the docket to read background documents or the received electronic and written/paper comments, go to https://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 Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT:
Theresa Wells, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 1, Rm. 1202, Silver Spring, MD 20993, 703–380–3900. Theresa.wells@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

Nearly all tobacco product use begins during youth and young adulthood (Ref. 1). While the current use of any tobacco product among U.S. middle and high school students has decreased from 2011–2017, there has been an alarming increase in e-cigarette use over this time. In fact, since 2014, e-cigarettes have been the most commonly used tobacco products among youth, used by 1.73 million (11.7 percent) high school students and 390,000 (3.3 percent) middle school students in 2017 (Ref. 2). Youth e-cigarette use raises a number of health concerns including risk of addiction to nicotine early on in life, potential harm to the developing adolescent brain, and exposure to chemicals including carbonyl compounds and volatile organic compounds known to have adverse health effects; the full range of possible health effects is not yet completely understood (Ref. 3).

On April 24, 2018, FDA announced its Youth Tobacco Prevention Plan. This plan focuses on three key strategies: Prevention of youth access to tobacco products, curbing the marketing of tobacco products aimed at youth, and educating teens about the dangers of using any tobacco products. FDA recently launched an expansion of its “The Real Cost” campaign to educate youth on the dangers of e-cigarette use and increased enforcement actions to

[1] An e-cigarette is one type of electronic nicotine delivery system, which also includes e-cigars, e-hooakah, vape personal vaporizers, and electronic pipes. See https://www.fda.gov/TobaccoProducts/Labeling/Products IngredientsComponents/ucm456610.htm and Ref. 2.

address this critically important public health concern. In addition to the prevention of initiation, which will be the cornerstone of any successful effort to curb youth e-cigarette use, FDA is also exploring additional approaches to address youth e-cigarette use. One such approach may be the development of drug therapies, as part of multimodal treatment strategies, including behavioral interventions, to support tobacco product cessation. To date, research on youth tobacco product cessation has been limited and focused on smoking (i.e., combustible products) cessation. One recent review found a paucity of data on either behavioral or drug therapies for smoking cessation in young people (age less than 20 years) and concluded that “there continues to be a need for well-designed, adequately powered, randomized controlled trials of interventions for this population of smokers” (Ref. 4). FDA is not aware of any research examining either drug or behavioral interventions for the cessation of youth or adult e-cigarette use. In contrast, there is a large body of research on adult smoking cessation, and multiple drugs for smoking cessation are approved for the adult population, including a variety of prescription and over-the-counter nicotine replacement therapy (NRT) products, as well as the prescription drugs varenicline and bupropion hydrochloride sustained release (see Appendix A).

II. Purpose and Scope of the Public Hearing

FDA is holding a public hearing to obtain the public’s perspectives on the potential role drug therapies may play in the broader effort to eliminate youth e-cigarette and other tobacco product use, as well as the appropriate methods and study designs for evaluating youth e-cigarette cessation therapies and the safety and efficacy of such therapies. The Agency has determined that a public hearing is the most appropriate way to ensure public engagement on this issue, which is of great importance to the public health. FDA believes it is critical to obtain input across the medical and research fields, the pharmaceutical and tobacco industries, and among public health stakeholders (including adolescents) regarding approaches to eliminate youth e-cigarette and other tobacco product use, including exploring whether there is a need for drug therapies to support youth e-cigarette cessation, and if so, how FDA

Questions for Commenters To Address

Considering the broad range of activities focused on this public health issue, FDA is interested in the public’s view on approaches to eliminating e-cigarette and other tobacco product use among youth. Although FDA welcomes all feedback on any public health, scientific, regulatory or legal considerations relating to this topic, we particularly encourage commenters to consider the following questions as they prepare their comments or statements. Responses to questions should include supporting scientific justification.

1. FDA notes that the factors driving e-cigarette use among youth likely differ from those in the adult population. How might such differences impact the need for, or use of, drug therapies for e-cigarette cessation among youth?

2. Is there evidence to suggest that drug therapies are appropriate for assessment in youth e-cigarette cessation? What age groups (older adolescent vs. younger adolescent), patterns in tobacco use (duration and frequency of use), and clinical features (level of addiction, presence/absence of comorbidities including psychiatric disease) might characterize this population? What types of products (NRT vs. non-NRT; prescription vs. over-the-counter) might be useful?

3. Describe the scientific, clinical, and societal factors that could either encourage or impede the conduct of clinical trials designed to evaluate drugs intended for youth e-cigarette cessation. What approaches could be used to encourage research and overcome barriers to research?

4. What methods and study designs are appropriate for assessing drug therapies for youth e-cigarette cessation? What are the appropriate control groups? What are the most informative endpoints and the best assessment tools to evaluate these endpoints?

5. Acknowledging that to date research has been limited, are there data available from the adult experience with smoking cessation that could potentially be leveraged in the effort to develop drug therapies for youth e-cigarette cessation? Have any drug therapies demonstrated potential to help adults discontinue e-cigarette use? Are there differences between adolescents and adults that impact the ability to extrapolate efficacy findings from the adult population to the adolescent population? Could existing NRT products be useful for youth e-cigarette cessation?

6. While this hearing is focused on the topic of e-cigarette use among youth, as e-cigarettes are currently the most commonly used form of tobacco in this population, FDA also welcomes comments regarding the potential need for drug therapies to support cessation of other tobacco products, including combustible products (i.e., cigarettes or cigars) and smokeless tobacco products, among youth and the issues impacting the development of such therapies.

Registration and Requests for Oral Presentations: The FDA Conference Center at the White Oak location is a Federal facility with security procedures and limited seating. Attendance will be free and on a first-come, first-served basis. For those interested in presenting at the meeting, either with a formal oral presentation or as a speaker in the open public hearing session, please register by Friday, November 23, 2018, at https://www.eventbrite.com/e/fda-pediatric-tobacco-cessation-part-15-public-hearing-tickets-50167147288. If you wish to attend either in person or by Webcast (see Streaming Webcast of the Public Hearing), please register for the hearing by Monday December 3, 2018, at https://www.eventbrite.com/e/fda-pediatric-tobacco-cessation-part-15-public-hearing-tickets-50167147288. Those without internet or email access can register and/or request to participate as an open public hearing speaker or a formal presenter by contacting Theresa Wells by the above dates (see FOR FURTHER INFORMATION CONTACT).

FDA will try to accommodate all persons who wish to make a presentation. Formal oral presenters may use an accompanying slide deck, while those participating in the Open Public Hearing will have less allotted time than formal oral presenters and will deliver oral testimony only (no accompanying slide deck). Individuals wishing to present should identify the number of the specific question, or questions, they wish to address. This will help FDA organize the presentations. Individuals and organizations with a financial interest should consolidate or coordinate their presentations and request time for a joint presentation. Individual organizations are limited to a single presentation slot. FDA will notify registered presenters of their scheduled presentation times. The time allotted for each presentation will depend on the number of individuals who wish to speak. Registered presenters making a formal oral presentation are encouraged to submit an electronic copy of their presentation (Powerpoint or PDF) to OMPTEFeedback@fda.hhs.gov with the subject line “Eliminating Youth Electronic Cigarette and Other Tobacco Product Use: The Role for Drug Therapies” on or before Wednesday, November 28, 2018. Persons registered to present are encouraged to arrive at the hearing room early and check in at the onsite registration table to confirm their designated presentation time. Actual presentation times, however, may vary based on how the meeting progresses in real-time. An agenda for the hearing and any other background materials will be made available 5 days before the hearing at https://www.fda.gov/NewsEvents/MeetingsConferencesWorkshops/ucm620744.htm.

If you need special accommodations because of a disability, please contact Theresa Wells (see FOR FURTHER INFORMATION CONTACT) at least 7 days before the hearing.

Streaming Webcast of the Public Hearing: For those unable to attend in person, FDA will provide a live Webcast of the hearing. To join the hearing via the Webcast, please go to https://collaboration.fda.gov/ptc120518.

Transcripts: Please be advised that as soon as a transcript is available, it will be accessible at http://www.regulations.gov. It may be viewed at the Dockets Management Staff (see Comments). A transcript will also be available in either hard copy or on CD-ROM, after submission of a Freedom of Information request. The Freedom of Information office address is available on the Agency’s website at https://www.fda.gov.

III. Notice of Hearing Under 21 CFR Part 15

The Commissioner of Food and Drugs is announcing that the public hearing will be held in accordance with 21 CFR part 15. The hearing will be conducted by a presiding officer, who will be accompanied by FDA senior management from the Office of the Commissioner, the Center for Drug Evaluation and Research, and the Center for Tobacco Products. Under § 15.30(f), the hearing is informal and the rules of evidence do not apply. No participant may interrupt the presentation of another participant. Only the presiding officer and panel members can pose questions; they can question any person during or at the conclusion of each presentation. Public hearings under part 15 are subject to FDA’s policy and procedures for electronic media coverage of FDA’s public administrative proceedings (21 CFR part 10, subpart C). Under § 10.205, representatives of the media may be permitted, subject to certain limitations, to videotape, film, or otherwise record FDA’s public
administrative proceedings, including presentations by participants. The hearing will be transcribed as stipulated in § 15.30(b) (see Transcripts). To the extent that the conditions for the hearing, as described in this notice, conflict with any provisions set out in part 15, this notice acts as a waiver of those provisions as specified in § 15.30(h).

IV. References

The following references marked with an asterisk (*) are on display at the Dockets Management Staff and are available for viewing by interested persons between 9 a.m. and 4 p.m., Monday through Friday; they also are available electronically at https://www.regulations.gov. References without asterisks are not on public display at https://www.regulations.gov because they have copyright restriction. Some may be available at the website address, if listed. References without asterisks are available for viewing only at the Dockets Management Staff. FDA has verified the website addresses, as of the date this document publishes in the Federal Register, but websites are subject to change over time.


BILLING CODE 4164–01–P

Appendix A: Summary of FDA-Approved Active NDAs of NRTs and non-NRTs Indicated for Smoking Cessation
(October 5, 2018)

<table>
<thead>
<tr>
<th>Product Name (NDA #, holder)</th>
<th>OTC or Rx (Date approved, Date Rx→OTC)</th>
<th>Route (Doses)</th>
<th>Indication</th>
<th>Adult Treatment Duration and Schedule</th>
<th>Pediatric Labeling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nicorette gum (nicotine polacrilex) (NDA 018612 for 2 mg, NDA 020666 for 4 mg, GSK)</td>
<td>Approved as prescription on 1/13/84 for 2 mg, 6/8/92 for 4 mg, Rx→OTC for both on 2/9/96.</td>
<td>Oral (2, 4 mg gum)</td>
<td>Reduces withdrawal symptoms, including nicotine craving, associated with quitting smoking.</td>
<td>12 weeks (for longer use, talk to healthcare provider):</td>
<td>If you are under 18 years of age ask a doctor before use.</td>
</tr>
</tbody>
</table>

If smoke 1st cigarette within 30 min of waking up, use 4 mg; if more than 30 min, use 2 mg. |

NRT Therapies

Nicoderm CQ (nicotine) (NDA 020616; Sanofi Aventis) | Approved as prescription on 11/7/91, Rx→OTC on 8/2/96. | Patch (7, 14, 21 mg) | Same as above | 10 weeks and 8 weeks (for longer use, talk to healthcare provider): | Same as above |

If ≥ 10 cigarettes/day: |

Wk 1: 6 mg/day |

Wk 7-8: 1 mg/day |

Wk 9-10: 6 mg/day |

If ≤ 10 cigarettes/day: |

Wk 1: 6 mg/day |

Wk 7-8: 1 mg/day |

Wk 9-10: 6 mg/day |

Habitrol (nicotine) (NDA 02078; Dr. Reddy’s) | Approved as prescription on 11/27/99, Rx→OTC on 11/12/99. | Patch (7, 14, 21 mg) | Same as above | 8 weeks (for longer use, talk to healthcare provider): | Same as above |

If ≥ 10 cigarettes/day: |

Wk 1: 1 mg/day |

Wk 5-6: 6 mg/day |

Wk 7-8: 1 mg/day |

If ≤ 10 cigarettes/day: |

Wk 1: 1 mg/day |

Wk 5-6: 6 mg/day |

Wk 7-8: 1 mg/day |
<table>
<thead>
<tr>
<th>Product Name (NDA #; holder)</th>
<th>OTC or Rx (Date approved, Date Rx→OTC)</th>
<th>Route (Doses)</th>
<th>Indication</th>
<th>Adult Treatment Duration and Schedule</th>
<th>Pediatric Labeling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nicotrol NS (nicotine) (NDA 020385; Pfizer)</td>
<td>Prescription (3/22/96; N/A)</td>
<td>Nasal spray (50 microliter spray delivering 0.5 mg)</td>
<td>• Indicated as an aid to smoking cessation for the relief of nicotine withdrawal symptoms • Should be used as a part of a comprehensive behavioral smoking cessation program</td>
<td>2 sprays (one per nostril) = 1 dose • Starting dose: 1-2 doses/hour • Maximum doses/hour: 5 • Maximum doses/day: 10</td>
<td>The safety and efficacy of the continued use of Nicotrol NS for periods longer than 6 months have not been adequately studied and such use is not recommended. Under Pediatric Use: Not recommended for use in the pediatric population because its safety and effectiveness in children and adolescents who smoke have not been evaluated.</td>
</tr>
<tr>
<td>Nicotrol Inhaler (nicotine) (NDA 020714; Pharmacia and Upjohn)</td>
<td>Prescription (5/2/97; N/A)</td>
<td>Inhalant (10 mg cartridge; 4 mg delivered)</td>
<td>• Indicated as an aid to smoking cessation for the relief of nicotine withdrawal symptoms • Recommended for use as part of a comprehensive behavioral smoking cessation program.</td>
<td>The recommended duration of treatment is 3 months, after which patients may be weaned from the inhaler by gradual reduction of the daily dose over the following 6 to 12 weeks. The safety and efficacy of the continued use of Nicotrol Inhaler for periods longer than 6 months have not been studied and such use is not recommended. Safety and effectiveness in pediatric and adolescent patients below the age of 18 years have not been established for any nicotine replacement product. However, no specific medical risk is known or expected in nicotine dependent adolescents. NICOTROL Inhaler should be used for the treatment of tobacco dependence in the older adolescent only if the potential benefit justifies the potential risk.</td>
<td></td>
</tr>
<tr>
<td>Nicorette lozenge (nicotine polacrilex) (NDA 021330; OSK)</td>
<td>OTC (10/31/02; N/A)</td>
<td>Oral (2, 4 mg)</td>
<td>Reduces withdrawal symptoms, including nicotine craving, associated with quitting smoking.</td>
<td>12 weeks (for longer use, talk to health care provider): • Wk 1-6: 1 per 1-2 hr • Wk 7-9: 1 per 2-4 hr • Wk 10-12: 1 per 4-8 hr If smoke 1st cigarette within 30 min of waking up, use 4 mg; if more than 30 min, use 2 mg.</td>
<td>If you are under 18 years of age ask a doctor before use. No studies have been done to show if this product will work for you.</td>
</tr>
<tr>
<td>Nicorette mini lozenge (nicotine polacrilex) (NDA 022360; OSK)</td>
<td>OTC (5/18/09; N/A)</td>
<td>Oral (2, 4 mg)</td>
<td>Same as above</td>
<td>Same as above</td>
<td>Same as above</td>
</tr>
</tbody>
</table>

**Non-NRT Therapies**

<table>
<thead>
<tr>
<th>Product Name (NDA #; holder)</th>
<th>OTC or Rx (Date approved, Date Rx→OTC)</th>
<th>Route (Doses)</th>
<th>Indication</th>
<th>Adult Treatment Duration and Schedule</th>
<th>Pediatric Labeling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zyban (bupropion hydrochloride sustained release) (NDA 020711; OSK)</td>
<td>Prescription (5/14/97; N/A)</td>
<td>Oral (150 mg)</td>
<td>• Indicated as an aid to smoking cessation treatment</td>
<td>7-12 weeks: • Start at one 150-mg tablet per day for 3 days • Can increase to 300 mg per day given as one 150-mg tablet twice each day, with 8 hours between • Patient may benefit from ongoing treatment.</td>
<td>Safety and effectiveness in the pediatric population have not been established. Boxed Warning for suicidality in children, adolescents, and young adults in setting of bupropion use as an antidepressant.</td>
</tr>
</tbody>
</table>

Leslie Kux, Associate Commissioner for Policy.

[FR Doc. 2018–24126 Filed 11–2–18; 8:45 am]
BILLING CODE 4164–01–C

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

21 CFR Part 101

[Docket No. FDA–2018–D–1459]

Food Labeling: Serving Sizes of Foods That Can Reasonably Be Consumed at One Eating Occasion, Reference Amounts Customarily Consumed, Serving Size-Related Issues, Dual-Column Labeling, and Miscellaneous Topics; Draft Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notification of availability.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of a draft guidance for industry entitled “Food Labeling: Serving Sizes of Foods That Can Reasonably Be Consumed at One Eating Occasion, Reference Amounts Customarily Consumed, Serving Size-Related Issues, Dual-Column Labeling, and Miscellaneous Topics.” The draft guidance, when finalized, will provide questions and answers on topics related primarily to implementing two final rules, one entitled “Food Labeling: Serving Sizes of Foods That Can Reasonably Be Consumed At One Eating Occasion; Reference Amounts Customarily Consumed, Serving Size-Related Issues, Dual-Column Labeling, and Miscellaneous Topics.” The draft guidance also discusses formatting issues for dual-column labeling, products that have limited space for nutrition labeling, and additional issues dealing with compliance.

DATES: Submit either electronic or written comments on the draft guidance by January 4, 2019 to ensure that we consider your comment on the draft guidance before we begin work on the final version of the guidance.

ADDRESSES: You may submit comments on any guidance at any time as follows:

Electronic Submissions

Submit electronic comments in the following way:

- Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://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 https://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): Dockets Management Staff (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

- For written/paper comments submitted to the Dockets Management Staff, 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–2018–D–1459 for “Food Labeling: Serving Sizes of Foods That Can Reasonably Be Consumed At One Eating Occasion, Reference Amounts Customarily Consumed, Serving Size-Related Issues, Dual-Column Labeling, and Miscellaneous Topics; Draft Guidance for Industry.” Received comments will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at https://www.regulations.gov or at the Dockets Management Staff 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.” We will review this copy, including the claimed confidential information, in our 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 https://www.regulations.gov. Submit both copies to the Dockets Management Staff. 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: https://www.gpo.gov/fdsys/pkg/FR-2015-09-18/pdf/2015-23389.pdf.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to https://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 Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

You may submit comments on any guidance at any time (see 21 CFR 10.115(g)(5)).

Submit written requests for single copies of the draft guidance to the Office of Nutrition and Food Labeling, Center for Food Safety and Applied Nutrition, Food and Drug Administration, 5001 Campus Dr., College Park, MD 20740. Send two self-addressed adhesive labels to assist that office in processing your request. See the SUPPLEMENTARY INFORMATION section for electronic access to the draft guidance.

FOR FURTHER INFORMATION CONTACT:
Jillonne Kevala, Center for Food Safety and Applied Nutrition, Food and Drug Administration, 5001 Campus Dr., College Park, MD 20740, 240–402–1450.

SUPPLEMENTARY INFORMATION:

I. Background

We are announcing the availability of a draft guidance for industry entitled “Food Labeling: Serving Sizes of Foods That Can Reasonably Be Consumed At One Eating Occasion, Reference Amounts Customarily Consumed, Serving Size-Related Issues, Dual-