Interested persons may observe the deliberations, but the Committee will not hear further comments during this time except at the request of the chairperson. The Committee will also allow a 15-minute unscheduled open public session for any attendee to address issues specific to the topics under consideration. At the conclusion of the day, the members will vote and the Committee will make its recommendation(s) to CMS.

III. Registration Instructions

CMS’ Coverage and Analysis Group is coordinating meeting registration. While there is no registration fee, individuals must register to attend. You may register online at http://www.cms.gov/apps/events/upcomingevents.asp?OrderBy=1&type=3 or by phone by contacting the person listed in the FOR FURTHER INFORMATION CONTACT section of this notice by the deadline listed in the DATES section of this notice. Please provide your full name (as it appears on your state-issued driver’s license), address, organization, telephone number(s), fax number, and email address. You will receive a registration confirmation with instructions for your arrival at the CMS complex or you will be notified that the seating capacity has been reached.

IV. Security, Building, and Parking Guidelines

This meeting will be held in a federal government building; therefore, federal security measures are applicable. The Real ID Act, enacted in 2005, establishes minimum standards for the issuance of state-issued driver’s licenses and identification (ID) cards. It prohibits Federal agencies from accepting an official driver’s license or ID card from a state unless the Department of Homeland Security determines that the state meets these standards. Beginning October 2015, photo IDs (such as a valid driver’s license) issued by a state or territory not in compliance with the Real ID Act will not be accepted as identification to enter Federal buildings. Visitors from these states/territories will need to provide alternative proof of identification (such as a valid passport) to gain entrance into CMS buildings.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Determination That SODIUM IODIDE I-123 (Sodium Iodide I-123), Oral Solution, 2 Millicuries/Milliliter, Was Not Withdrawn From Sale for Reasons of Safety or Effectiveness

AGENCY: Food and Drug Administration, HHS.
Silver Spring, MD 20993–0002, 301–977–8900. Answers to commonly asked questions including information regarding special accommodations due to a disability, visitor parking, and transportation may be accessed at: https://www.fda.gov/AdvisoryCommittees/AboutAdvisoryCommittees/ucm408555.htm.

FOR FURTHER INFORMATION CONTACT:
Evela Washington, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, Rm. G640, Silver Spring, MD 20933–0002, 301–796–6683, or FDA

Department of Health and Human Services
Food and Drug Administration
[Docket No. FDA–2018–N–1726]
Circulatory System Devices Panel of the Medical Devices Advisory Committee; Notice of Meeting

A person may petition the Agency to determine, or the Agency may determine on its own initiative, whether a listed drug was withdrawn from sale for reasons of safety or effectiveness. This determination will allow FDA to approve abbreviated new drug applications (ANDAs) for a previously approved abbreviated new drug application (ANDA) that does not refer to a listed drug.—SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, was withdrawn from sale for reasons of safety or effectiveness. This determination may be made at any time after the drug has been withdrawn from sale. It must be made prior to FDA’s approval of an ANDA that refers to the listed drug (§ 314.161). FDA may not approve an ANDA that does not refer to a listed drug.

FDA’s approval of an ANDA that refers to a listed drug may be made at any time after the drug has been withdrawn from sale. It must be made prior to FDA’s approval of an ANDA that refers to the listed drug (§ 314.161). FDA may not approve an ANDA that does not refer to a listed drug.

SUMMARY:
ACTION:
The Food and Drug Administration (FDA or Agency) has determined that SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, was not withdrawn from sale for reasons of safety or effectiveness. This determination will allow FDA to approve abbreviated new drug applications (ANDAs) for sodium iodide I 123, oral solution, 2 mCi/mL, if all other legal and regulatory requirements are met.

FOR FURTHER INFORMATION CONTACT:

SUPPLEMENTARY INFORMATION:
In 1984, Congress enacted the Drug Price Competition and Patent Term Restoration Act of 1984 (Pub. L. 98–417) (the 1984 amendments), which authorized the approval of duplicate versions of drug products under an ANDA procedure. ANDA applicants must, with certain exceptions, show that the drug for which they are seeking approval contains the same active ingredient in the same strength and dosage form as the “listed drug,” which is a version of the drug that was previously approved. ANDA applicants do not have to repeat the extensive clinical testing otherwise necessary to gain approval of a new drug application (NDA).

The 1984 amendments include what is now section 505(j)(7) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 355(j)(7)), which requires FDA to publish a list of all approved drugs. FDA publishes this list as part of the “Approved Drug Products With Therapeutic Equivalence Evaluations,” which is known generally as the “Orange Book.” Under FDA regulations, drugs are removed from the list if the Agency withdraws or suspension approval of the drug’s NDA or ANDA for reasons of safety or effectiveness or if FDA determines that the listed drug was withdrawn from sale for reasons of safety or effectiveness (§ 314.162 (21 CFR 314.162)).

A person may petition the Agency to determine, or the Agency may determine on its own initiative, whether a listed drug was withdrawn from sale for reasons of safety or effectiveness. This determination may be made at any time after the drug has been withdrawn from sale. It must be made prior to FDA’s approval of an ANDA that refers to the listed drug (§ 314.161 (21 CFR 314.161)). FDA may not approve an ANDA that does not refer to a listed drug.

SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, is the subject of NDA 017630, held by GE Healthcare, and initially approved on March 24, 1976. SODIUM IODIDE I 123 is indicated for use in the evaluation of thyroid function or morphology.

SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, was withdrawn from sale for reasons of safety or effectiveness.

After considering the citizen petition and reviewing Agency records and based on the information we have at this time, FDA has determined under § 314.161 that SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, was withdrawn from sale for reasons of safety or effectiveness. The petitioner has identified no data or other information suggesting that SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, was withdrawn from sale for reasons of safety or effectiveness. We have carefully reviewed our files for records concerning the withdrawal of SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, from sale. We have also independently evaluated relevant literature and data for possible postmarketing adverse events. We have found no information that would indicate that this drug product was withdrawn from sale for reasons of safety or effectiveness.

Accordingly, the Agency will continue to treat SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, in the “Discontinued Drug Product List” section of the Orange Book. The “Discontinued Drug Product List” delineates, among other items, drug products that have been discontinued from marketing for reasons other than safety or effectiveness. ANDAs that refer to SODIUM IODIDE I 123 (sodium iodide I-123), oral solution, 2 mCi/mL, may be approved by the Agency as long as they meet all other legal and regulatory requirements for the approval of ANDAs. If FDA determines that labeling for this drug product should be revised to meet current standards, the Agency will advise ANDA applicants to submit such labeling.

Dated: May 7, 2018.

Leslie Kux,
Associate Commissioner for Policy.

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration
[Docket No. FDA–2018–N–1726]
Circulatory System Devices Panel of the Medical Devices Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) announces a forthcoming public advisory committee meeting of the Circulatory System Devices Panel of the Medical Devices Advisory Committee. The general function of the committee is to provide advice and recommendations to the Agency on FDA’s regulatory issues. The meeting will be open to the public.

DATES: The meeting will be held on June 12, 2018, from 8 a.m. to 6 p.m.