which is submitted annually by plans. Additionally, the proposed collection will request information to be split out by the authority under which each plan offers the benefits (mandatory, optional, mandatory-SSBCI, mandatory-Uniformity Flexibility). Form Number: CMS–10261 (OMB control number: 0938–1054); Frequency: Annually; Affected Public: Business or other for-profits; Number of Respondents: 743; Total Annual Responses: 6,687; Total Annual Hours: 187,979. (For policy questions regarding this collection contact Lucia Patrone at (410) 786–8621).


William N. Parham, III, Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2023–20739 Filed 9–22–23; 8:45 am]
BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2023–N–2079]

Hospira, Inc., et al.; Withdrawal of Approval of Eight Abbreviated New Drug Applications; Correction

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice; correction.

SUMMARY: The Food and Drug Administration (FDA) is correcting a notice entitled “Hospira, Inc., et al.; Withdrawal of Approval of Eight Abbreviated New Drug Applications” that appeared in the Federal Register of June 2, 2023. The document announced the withdrawal of approval (as of July 3, 2023) of eight abbreviated new drug applications (ANDAs) from multiple applicants. The document indicated that FDA was withdrawing approval of ANDA 077029, Calcipotriene Solution, 0.005% after receiving a withdrawal request from Tolmar, Inc., 701 Centre Ave., Fort Collins, CO 80526. Before FDA withdrew the approval of this ANDA, Tolmar, Inc. informed FDA that it did not want the approval of the ANDA withdrawn. Because Tolmar, Inc. timely requested that approval of ANDA 077029 not be withdrawn, the approval is still in effect. This document corrects that error.

FOR FURTHER INFORMATION CONTACT: Martha Nguyen, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 75, Rm. 1676, Silver Spring, MD 20993–0002, 240–402–6980, Martha.Nguyen@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In the Federal Register of Friday, June 2, 2023 (88 FR 36320), in FR Doc. 2023–11744, the following correction is made:

On page 36321, in the table, the entry for ANDA 077029 is removed.


Lauren K. Roth, Associate Commissioner for Policy.

[FR Doc. 2023–20742 Filed 9–22–23; 8:45 am]
BILLING CODE 4164–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Yoga Products Deviating From Standard of Identity; Amendment of Temporary Marketing Permit]

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or we) are amending the standard of identity for yogurt with modified ingredients, and yogurt deviating from standard of identity for yogurt with modified ingredients, and yogurt deviating from lower-fat yogurt products deviating from the requirements for the basic dairy ingredient provision of the yogurt standard of identity under 21 CFR 131.200(b).

Under our regulations at 21 CFR 130.17(f), we are amending the temporary permit issued to Chobani, LLC, to allow the test product to be used as ingredients, in whole or in part, in other nonstandardized foods. All other conditions and terms of this permit remain the same.


Lauren K. Roth, Associate Commissioner for Policy.

[FR Doc. 2023–20736 Filed 9–22–23; 8:45 am]
BILLING CODE 4164–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Heart, Lung, and Blood Institute; Amended Notice of Meeting

Notice is hereby given of a change in the meeting of the National Heart, Lung, and Blood Institute Initial Review Group, Mentored Patient-Oriented Research Study Section, October 26, 2023, 10:00 a.m. to October 27, 2023, 6 p.m., National Institutes of Health, Rockledge 1, 6705 Rockledge Drive, Bethesda, MD 20892, which was published in the Federal Register on September 14, 2023, FR Doc 2023–19907, 88 FRN 63115.

The National Heart, Lung, and Blood Institute Initial Review Group, Mentored Patient-Oriented Research Study Section meeting is being amended due to a change of the meeting time formats. The meeting will be held on October 26, 2023, from 9:00 a.m. to October 27, 6:00 p.m. This meeting will be a video assisted meeting, and closed to the public.

Dated: September 18, 2023.

Melanie J. Pantoya, Program Analyst, Office of Federal Advisory Committee Policy.

[FR Doc. 2023–20648 Filed 9–22–23; 8:45 am]
BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Human Genome Research Institute; Notice of Closed Meetings

Pursuant to section 1009 of the Federal Advisory Committee Act, as