SUPPLEMENTARY INFORMATION section for registration date and information.

ADDRESSES: Please note that due to the impact of the COVID–19 pandemic, all participants will be joining this public workshop via an online teleconferencing platform.

FOR FURTHER INFORMATION CONTACT:

Rokhsareh Shahidzadeh, Office of Scientific Professional Development, Office of the Chief Scientist, Office of the Commissioner, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 32, Rm. 2383, Silver Spring, MD 20993, 301–796–8740, *FDASciProDev@fda.hhs.gov.*

SUPPLEMENTARY INFORMATION:

I. Background

The FDA Science Forum is held biennially to inform the public about the groundbreaking science conducted at the Agency and to show how scientific research is used in FDA's regulatory decisions to protect and promote public health. Open to the public, industry, academia, patient advocates, government agencies, and current and potential collaborators, the 2-day event offers an opportunity to hear FDA scientific experts and nationally renowned scientists speak on a range of topics associated with regulatory science.

II. Topics for Discussion at the Public Workshop

The theme for the 2021 FDA Science Forum, "Science as the Foundation for Protecting and Promoting Public Health", will highlight areas of FDA research, including: (1) Improving clinical and postmarket evaluation; (2) substance use, misuse, and addiction; (3) product development and manufacturing; and (4) medical countermeasures, infectious disease, and pathogen-reduction technologies.

III. Participating in the Public Workshop

Registration: To register for the public workshop, please visit the following website: https://www.fda.gov/ scienceforum. Registration is free. Persons interested in attending this public workshop must register by May 21, 2021, by 5 p.m. Eastern Time. Registrants will receive confirmation when they have been accepted.

If you need special accommodations due to a disability, please contact Rokhsareh Shahidzadeh (see FOR FURTHER INFORMATION CONTACT) no later than May 21, 2021, by 5 p.m. Eastern Time.

Streaming Webcast of the Public Workshop: This public workshop will be streamed via a webcast only. To register for the webcast, please visit the following website: *https://www.fda.gov/ scienceforum.* Participants interested in viewing via webcast must register by May 21, 2021, by 5 p.m. Eastern Time.

If you have never attended a Connect Pro event before, test your connection at https://collaboration.fda.gov/common/ help/en/support/meeting_test.htm. To get a quick overview of the Connect Pro program, visit https://www.adobe.com/ go/connectpro_overview. FDA has verified the website addresses in this document, as of the date this document publishes in the **Federal Register**, but websites are subject to change over time.

Dated: March 24, 2021.

Lauren K. Roth,

Acting Principal Associate Commissioner for Policy.

[FR Doc. 2021–06705 Filed 3–31–21; 8:45 am] BILLING CODE 4164–01–P

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2020-N-2306]

TG United, Inc., et al.; Withdrawal of Approval of 27 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 that appeared in the Federal Register of January 15, 2021. The document announced the withdrawal of approval of 27 abbreviated new drug applications (ANDAs) from multiple applicants as of February 16, 2021. The document indicated that FDA was withdrawing approval of the following three ANDAs after receiving a withdrawal request from Upsher-Smith Laboratories, LLC., 6701 Evenstad Dr., Maple Grove, MN 55369: ANDA 084041, Chlordiazepoxide Hydrochloride (HCl) Capsules, 10 milligrams (mg); ANDA 084678, Chlordiazepoxide HCl Capsules, 5 mg; and ANDA 084679, Chlordiazepoxide HCl Capsules, 25 mg. Before FDA withdrew the approval of these ANDAs, Upsher-Smith Laboratories, LLC., informed FDA that it did not want the approval of the ANDAs withdrawn. Because Upsher-Smith Laboratories, LLC., timely requested that approval of these ANDAs not be withdrawn, the approval of ANDAs 084041, 084678, and 084679 is still in effect. In addition, the document indicated that FDA was

withdrawing approval of ANDA 206061, Pravastatin Sodium Tablets, 20 mg, 40 mg, and 80 mg, after receiving a request from Hisun Pharmaceutical (Hangzhou) Co., Ltd. However, the document published with the incorrect applicant name for ANDA 206061. This document corrects that error. All other information for ANDA 206061 remains the same.

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, January 15, 2021 (86 FR 4081), appearing on page 4081 in FR Doc. 2021–00833, the following corrections are made on page 4082 in the table:

1. The entries for ANDAs 084041, 084678, and 084679 are removed.

2. In the third column, third item from the bottom, the applicant name "Hisun Pharmaceuticals USA, Inc." is corrected to read "Hisun Pharmaceuticals USA, Inc., U.S. Agent for Hisun Pharmaceutical (Hangzhou) Co., Ltd." for ANDA 206061.

Dated: March 29, 2021.

Lauren K. Roth,

Acting Principal Associate Commissioner for Policy.

[FR Doc. 2021–06745 Filed 3–31–21; 8:45 am] BILLING CODE 4164–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2021-N-0270]

Endocrinologic and Metabolic Drugs Advisory Committee; Notice of Meeting; Establishment of a Public Docket; Request for Comments

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice; establishment of a public docket; request for comments.

SUMMARY: The Food and Drug Administration (FDA) announces a forthcoming public advisory committee meeting of the Endocrinologic and Metabolic Drugs Advisory Committee. The general function of the committee is to provide advice and recommendations to FDA on regulatory issues. The meeting will be open to the public. FDA is establishing a docket for public comment on this document.