While existing guidances provide general principles and recommendations regarding comparability studies and management of manufacturing changes for biological products, they generally do not address specific CGT product challenges. The purpose of this draft guidance is to provide FDA's current thinking on: (1) management and reporting of manufacturing changes for CGT products based on a life-cycle approach and (2) comparability studies to assess the effect of manufacturing changes on CGT product quality.

This draft guidance is being issued consistent with FDA's good guidance practices regulation (21 CFR 10.115). The draft guidance, when finalized, will represent the current thinking of FDA on "Manufacturing Changes and Comparability for Human Cellular and Gene Therapy Products." It does not establish any rights for any person and is not binding on FDA or the public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations.

II. Paperwork Reduction Act of 1995

While this guidance contains no collection of information, it does refer to previously approved FDA collections of information. Therefore, clearance by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (PRA) (44 U.S.C. 3501-3521) is not required for this guidance. The previously approved collections of information are subject to review by OMB under the PRA. The collections of information in 21 CFR part 211 have been approved under OMB control number 0910-0139; the collections of information in 21 CFR part 312 have been approved under OMB control number 0910-0014; and the collections of information in 21 CFR 601.2 and 601.12 have been approved under OMB control number 0910-0338.

III. Electronic Access

Persons with access to the internet may obtain the draft guidance at https://www.fda.gov/vaccines-blood-biologics/guidance-compliance-regulatory-information-biologics/biologics-guidances, https://www.fda.gov/regulatory-information/search-fda-guidance-documents, or https://www.regulations.gov.

Dated: July 10, 2023.

Lauren K. Roth,

 $Associate\ Commissioner\ for\ Policy.$ [FR Doc. 2023–14917 Filed 7–13–23; 8:45 am]

BILLING CODE 4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2017-N-6395]

Request for Applications for New Members of the Clinical Trials Transformation Initiative/Food and Drug Administration Patient Engagement Collaborative

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice; request for applications.

SUMMARY: The Food and Drug Administration (FDA or Agency), in collaboration with the Clinical Trials Transformation Initiative (CTTI), is requesting applications from patient advocates interested in participating on the Patient Engagement Collaborative (PEC). The PEC is an ongoing, collaborative forum coordinated through the FDA's Patient Affairs Staff, Office of Clinical Policy and Programs (OCPP), Office of the Commissioner at FDA, and is hosted by CTTI. Through the PEC, the patient community and FDA Staff are able to discuss an array of topics related to increasing meaningful patient engagement with diverse populations in medical product development and regulatory discussions at FDA. The activities of the PEC may include, but are not limited to, providing diverse perspectives on topics such as systematic patient engagement, transparency, and communication; providing considerations for implementing new strategies to enhance patient engagement at FDA; and proposing new models of collaboration in which patient, caregiver and patient advocate perspectives are incorporated into general medical product development and regulatory processes. DATES: Applications can be submitted starting at 11:59 p.m. Eastern Time on July 14, 2023. This announcement is open to receive a maximum of 75 applications. Applications will be

happens first.

ADDRESSES: All applications should be submitted to FDA's Patient Affairs Staff in OCPP. The preferred application method is via the online submission system provided by CTTI, available at https://duke.qualtrics.com/jfe/form/SV_6L817z4YfyCHFVY. For those applicants unable to submit an application electronically, please call FDA's Patient Affairs Staff at 301–796–8460 to arrange for mail or delivery service submission.

accepted until 11:59 p.m. Eastern Time

applications are received, whichever

on August 14, 2023 or until 75

Only complete applications, as described under section IV of this document, will be considered.

FOR FURTHER INFORMATION CONTACT:

Wendy Slavit, Office of the Commissioner, Office of Clinical Policy and Programs, Patient Affairs Staff, Food and Drug Administration, 301– 796–8460,

PatientEngagementCollaborative@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background and Purpose

The CTTI is a public-private partnership cofounded by FDA and Duke University whose mission is to develop and drive adoption of practices that will increase the quality and efficiency of clinical trials. FDA and CTTI have long involved patients and considered patient perspectives in their work. Furthering the engagement of diverse patients as valued partners across the medical product research and development continuum requires an open forum for patients and regulators to discuss and exchange ideas.

The PEC is an ongoing, collaborative forum in which the patient community and FDA Staff discuss an array of topics related to increasing patient engagement in medical product development and regulatory discussions at FDA. The PEC is a joint endeavor between FDA and CTTI. The activities of the PEC may inform relevant FDA and CTTI activities. The PEC is not intended to advise or otherwise direct the activities of either organization, and membership will not constitute employment by either organization.

The Food and Drug Administration Safety and Innovation Act (Pub. L. 112-144), section 1137, entitled "Patient Participation in Medical Product Discussions," added section 569C to the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360bbb-8c). This provision directs the Secretary of Health and Human Services to "develop and implement strategies to solicit the views of patients during the medical product development process and consider the perspectives of patients during regulatory discussions." On November 4, 2014, FDA issued a Federal Register notice establishing a docket (FDA-2014–N–1698) for public commenters to submit information related to FDA's implementation of this provision. Upon review of the comments received, one common theme, among others, included establishing an external group to provide input on patient engagement strategies across FDA's Centers. After considering the comments, FDA formed the PEC in 2018 to discuss a variety of

patient engagement topics. This group is consistent with additional legislation subsequently enacted in section 3001 of the 21st Century Cures Act (Pub. L. 114–255) and section 605 of the FDA Reauthorization Act of 2017 (Pub. L. 115–52), further supporting tools for fostering patient participation in the

regulatory process.

The PEC currently has 16 members.
To help ensure continuity in its activities and organizational knowledge, the PEC maintains staggered membership terms. During the fall of 2023, eight members will complete a term and up to eight new members will be selected. The purpose of this notice is to announce that the application process for up to eight new members of the PEC is now open, and to invite and encourage applications by the submission deadline for appropriately qualified individuals.

II. Criteria for Membership

The PEC includes up to 16 diverse representatives of the patient community. Eight members from the previous application process will remain on the PEC. The current application process is to select up to eight new PEC members. Selected members will include the following: (1) patients who have personal experience with a disease or medical condition; (2) caregivers who help support a patientparent, child, partner, other family member, or friend—as they manage their disease or medical condition; and/ or (3) representatives of patient groups who, through their role in the patient group, have direct or indirect disease experience. Please note that for purposes of this activity, the term "caregiver" is not intended to include individuals who are engaged in caregiving as healthcare professionals; and the term "patient group" is used herein to encompass patient advocacy organizations, disease advocacy organizations, voluntary health agencies, nonprofit research foundations, and public health organizations. The ultimate goal of the application and selection process is to identify individuals who can represent patient voices for their patient community.

Selection criteria include the applicant's potential to meaningfully contribute to the activities of the PEC, ability to represent and express patient voices for their constituency, ability to work in a constructive manner with involved stakeholders, and understanding of the clinical research enterprise. Consideration will also be given to ensuring the PEC includes diverse perspectives and experiences,

including but not limited to sociodemographic factors (such as age, gender, ethnicity, and education level) and disease experience. PEC members are required to be residents of the United States and must be 18 years of age or older.

Financial and other conflicts of interest will not necessarily make applicants ineligible for membership in the PEC. However, applicants cannot be direct employees of the medical product development industry or a currently registered lobbyist for an FDA-regulated industry.

III. Responsibilities and Expectations

Participation as a PEC member is voluntary. Meetings will be held two to four times per year and will be conducted virtually with the potential for in-person events (in the Washington, DC area).

Reasonable accommodations will be made for members with special needs for participation in a meeting or for any necessary travel. Applications for PEC membership are encouraged from individuals of all ages, sexes, genders, sexual orientations, racial and ethnic groups, education levels, income levels, and those with and without disabilities. Travel support will be provided, as applicable.

To help ensure continuity in its activities and organizational knowledge, the PEC will maintain staggered membership terms for patient community representatives.

Membership terms for new members will be 2-year appointments, beginning January 1, 2024.

Additional responsibilities and expectations are set forth in the PEC Framework, which should be reviewed prior to submitting an application, and is available at https://ctti-clinicaltrials.org/wp-content/uploads/2023/05/PEC-Framework_Revised-Apr-10-2023 FINAL.pdf.

IV. Application Process

Any interested person may apply for membership on the PEC. To apply, go to https://duke.qualtrics.com/jfe/form/SV_6L8l7z4YfyCHFVY. The application is completed online and includes questions to help determine eligibility for the PEC, demographic and other background questions, and four brief essay questions. The brief essay questions, to be answered in 500 characters or fewer (including spaces), are as follows:

• Please explain why you would have an outstanding ability to represent and express the patient voice for the disease area(s) you selected above. • Please give a few examples of experiences that demonstrate your outstanding ability to work across or interact with stakeholders in the medical product development and regulatory processes.

 Please explain how you have established an understanding of the medical product development and

regulatory processes.

 Please tell us why you are interested in becoming a member of the PEC and how you would be able to contribute.

Completing the application also involves submitting: (1) A current onepage résumé or bio that summarizes your patient advocacy experience and related activities (PDF format required) and (2) a one-page letter of endorsement from a patient group (or other similar group) with which the applicant has worked closely on activities that are relevant to the PEC (PDF format required). Please note, only the application and the two documents specified above will be reviewed. Your completed application form, résumé or bio, and letter of endorsement should all be submitted at the same time.

The résumé or bio must provide examples and descriptions of relevant activities and experiences related to the applicant's qualifications for PEC membership. The letter of endorsement should emphasize information relevant to the criteria for membership described above. This letter must be from and written by someone other than yourself. The letter may address topics such as the applicant's involvement in patient advocacy activities, experiences that stimulated an interest in participating in discussions about patient engagement in medical product development and regulatory decision processes, and other information that may be helpful in evaluating the applicant's qualifications as a potential member of the PEC.

Applications will be accepted until 11:59 p.m. Eastern Time on August 14, 2023 or until 75 applications are received, whichever happens first. Only complete applications will be considered.

The application review period will take a minimum of 2 months after 11:59 p.m. Eastern Time on August 14, 2023.

Additional information may be needed from some applicants during the review period, including information relevant to understanding potential sources of conflict of interest, in which case applicants will be contacted directly. All applicants (both those selected for PEC membership and those who are not selected) will be notified of the final application decision no later than December 31, 2023.

Dated: July 10, 2023.

Lauren K. Roth,

Associate Commissioner for Policy. [FR Doc. 2023–14920 Filed 7–13–23; 8:45 am]

BILLING CODE 4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2023-D-0559]

Postmarketing Studies and Clinical Trials: Determining Good Cause for Noncompliance With Section 505(o)(3)(E)(ii) of the Federal Food, Drug, and Cosmetic Act; Draft Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of a draft guidance for industry entitled "Postmarketing Studies and Clinical Trials: Determining Good Cause for Noncompliance with Section 505(o)(3)(E)(ii) of the Federal Food, Drug, and Cosmetic Act." The Federal Food, Drug, and Cosmetic Act (FD&C Act) authorizes FDA to require certain postmarketing studies and clinical trials for prescription drugs at the time of approval or after approval if FDA becomes aware of new safety information. This draft guidance describes the factors FDA considers when determining whether an applicant has demonstrated good cause for failure to comply with the timetable for completion of studies or clinical trials required under the provisions. This draft guidance also provides information on relevant procedures, including how an applicant should communicate with FDA regarding compliance with these required studies and trials and describes actions FDA may take for noncompliance with the requirements.

DATES: Submit either electronic or written comments on the draft guidance by September 12, 2023 to ensure that the Agency considers your comment on this draft guidance before it begins 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–2023–D–0559 for "Postmarketing Studies and Clinical Trials: Determining Good Cause for Noncompliance with Section 505(o)(3)(E)(ii) of the Federal Food, Drug, and Cosmetic Act." 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, 240–402–7500.

• 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 dockets, see 80 FR 56469, September 18, 2015, or access the information at: https:// www.govinfo.gov/content/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, 240–402–7500.

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 Division of Drug Information, Center for Drug Evaluation and Research, Food and Drug Administration, 10001 New Hampshire Ave., Hillandale Building, 4th Floor, Silver Spring, MD 20993-0002; or Office of Communication, Outreach and Development, Center for Biologics Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 71, Rm. 3128, Silver Spring, MD 20993-0002. Send one self-addressed adhesive label to assist that office in processing your requests. See the SUPPLEMENTARY **INFORMATION** section for electronic access to the draft guidance document.

FOR FURTHER INFORMATION CONTACT:

Kathy Weil, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 22, Rm. 5367, Silver Spring, MD 20993, 301–796–6054, or Diane Maloney, Center for Biologics Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 71, Rm. 7301, Silver Spring, MD 20993–0002, 240–402–7911.