those strategies. The interview will focus on systems and processes and should not require preparation time by the Champion. The estimated time for the interview is two hours, which includes time to review the interview protocol with the interviewer, respond to the interview questions, and review a summary data about the Champion’s practices. The summary will be written as a success story and will be posted on the Million Hearts® Web site.

Additional Information: Information received from nominees will be stored in a password protected file on a secure server. The challenge Web site may post the number of nominations received but will not include confidential or proprietary information about individual nominees, as described further below. The database of information submitted by nominees will not be posted on the Web site. Information collected from nominees will include general details, such as the business name, address, and contact information of the nominee. This type of information is generally publicly available. The nomination will collect and store only aggregate clinical data through the nomination process; no individual identifiable patient data will be collected or stored. Confidential or propriety data, clearly marked as such, will be secured to the full extent allowable by law.

Information for selected Champions, such as the provider, practice, or health system’s name, location, hypertension control rate, and clinic practices that support hypertension control will be shared through press releases, the challenge Web site, and Million Hearts® and HHS/CDC resources.

Summary data on the types of systems and processes that all nominees use to control hypertension may be shared in documents or other communication products that describe generally used practices for successful hypertension control. HHS/CDC will use the summary data only as described. Compliance with Rules and Contacting Contest Winners: Finalists and Champions must comply with all terms and conditions of these Official Rules, and winning is contingent upon fulfilling all requirements herein. The initial finalists will be notified by email, telephone, or mail after the date of the judging.

Privacy: If Contestants choose to provide HHS/ CDC with personal information by registering or filling out the submission form through the Challenge.gov Web site, that information is used to respond to Contestants in matters regarding their submission, announcements of entrants, finalists, and winners of the Contest. Information is not collected for commercial marketing. Champions are permitted to cite that they were selected as Champions for the 2017 Million Hearts Hypertension Control Challenge. General Conditions: HHS/CDC reserves the right to cancel, suspend, and/or modify the Challenge, or any part of it, for any reason, at HHS/CDC’s sole discretion. If the Challenge is cancelled, suspended, and/or modified, HHS/CDC will inform the public through the publication of a notice in the Federal Register.

Participation in this Contest constitutes a contestants’ full and unconditional agreement to abide by the Contest’s Official Rules found at www.Challenge.gov.


Sandra Cashman,
Executive Secretary, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration

[Docket No. FDA–2017–N–0726]

Antibody Mediated Rejection in Kidney Transplantation; Public Workshop; Request for Comments

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of public workshop; request for comments.

SUMMARY: The Food and Drug Administration (FDA, the Agency, or we) is announcing a public workshop regarding new developments and scientific issues related to antibody mediated rejection (AMR) in kidney transplantation. This public workshop is intended to provide information for gain perspective from individuals, industry, health care professionals, researchers, public health organizations, patients, patient care providers, and other interested persons on various aspects of clinical development of medical products for prophylaxis and/or treatment of AMR in kidney transplant recipients, including clinical trial design and endpoints. The input from this public workshop will also help in developing topics for future discussion.

DATES: The public workshop will be held on April 12, 2017, from 8 a.m. to 6 p.m. and April 13, 2017, from 8:30 a.m. to 1:30 p.m. Submit either electronic or written comments on this public workshop by April 27, 2017.

Late, untimely filed comments will not be considered. Electronic comments must be submitted on or before April 27, 2017. The https://www.regulations.gov electronic filing system will accept comments until midnight Eastern Time at the end of April 27, 2017. 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. See the SUPPLEMENTARY INFORMATION section for registration date and information. Workshop updates and the workshop agenda will be made available at: http://www.fda.gov/Drugs/NewsEvents/ucm532070 prior to the workshop.

ADDRESSES: The public workshop will be held at the Tommy Douglas Conference Center, 10000 New Hampshire Ave., Silver Spring, MD 20903. The conference center’s phone number is 240–645–4000.

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 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): Division of Dockets Management (HFA–305), Food
and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.
• For written/paper comments submitted to the Division of Dockets Management, 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–2017–N–0726 for “Antibody Mediated Rejection in Kidney Transplantation.” Received comments, those filed in a timely manner (see DATES) will be placed in the docket and, except for information submitted, marked and identified, as confidential, except for information submitted, marked and identified, as confidential, if submitted as detailed in “Instructions.”

SUPPLEMENTARY INFORMATION:
I. Background
FDA is announcing a public workshop regarding AMR in kidney transplantation. This public workshop will focus on scientific considerations in the clinical development of medical products for prophylaxis and/or treatment of AMR in kidney transplant recipients.

Among the primary goals of this workshop are the discussion of the role of immunosuppressive medication nonadherence in the development of de novo donor specific antibody (DSA) formation and subsequent AMR, new developments in transplantation and their impact on patient management (such as pretransplant sensitization not manifested by DSA, donor/recipient human leukocyte antigen (HLA) epitope matching, routine posttransplant DSA monitoring), the natural course of the acute-chronic AMR continuum and its temporal association with cellular rejection and changes in glomerular filtration rate (GFR), unmet medical needs and the potential implications of these factors on the design of clinical trials for the prevention and management of AMR.

The Agency encourages individuals, industry, health care professionals, researchers, public health organizations, patients, patient care providers, and other interested persons to attend this public workshop.

II. Participating in the Public Workshop
Registration: Persons interested in attending this public workshop must register by April 6, 2017, midnight Eastern Time. Please provide complete contact information for each attendee, including name, title, affiliation, address, email, and telephone to AntibodyMediatedRejectionWorkshop2017@fda.hhs.gov no later than April 10, 2017. No commercial or promotional material will be permitted to be presented or distributed at the public workshop.

Requests for Oral Presentations: During online registration you may indicate if you wish to present during a public comment session or participate in a specific session, and which topic(s) you wish to address. We will do our best to accommodate requests to make public comments and requests to participate in the focused sessions. Individuals and organizations with common interests are urged to consolidate or coordinate their presentations, and request time for a joint presentation, or submit requests for designated representatives to participate in the focused sessions. All requests to make oral presentations must be received by the close of registration on April 6, 2017. Following the close of registration, we will determine the amount of time allotted to each presenter and the approximate time each oral presentation is to begin, and will select and notify participants by April 7, 2017. If selected for presentation, any presentation materials must be emailed to AntibodyMediatedRejectionWorkshop2017@fda.hhs.gov no later than April 10, 2017. No commercial or promotional material will be permitted to be presented or distributed at the public workshop.

Transcripts: Please be advised that as soon as a transcript of the public workshop is available, it will be accessible at https://www.regulations.gov. It may be viewed at the Division of Dockets Management (see ADDRESSES). A link to the transcript will also be available on the Internet at http://www.fda.gov/Drugs/NewsEvents/ucm532070.htm approximately 45 days after the workshop.


Anna K. Abram,
Deputy Commissioner for Policy, Planning, Legislation, and Analysis.

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