Early Childhood Home Visiting (MIECHV) Program when the participating children are in kindergarten. This Federal Register notice is seeking to extend data collection for the kindergarten followup. The original Federal Register notices for the MIHOPE-K data collection were titled under MIHOPE-Long-Term Follow-Up (MIHOPE-LT).

DATES: Comments due within 60 days of publication. In compliance with the requirements of the Paperwork Reduction Act of 1995, ACF is soliciting public comment on the specific aspects of the information collection described

ADDRESSES: Copies of the proposed collection of information can be obtained and comments may be forwarded by emailing OPREinfocollection@acf.hhs.gov. Alternatively, copies can also be obtained by writing to the Administration for Children and

Families, Office of Planning, Research, and Evaluation, 330 C Street SW, Washington, DC 20201, Attn: OPRE Reports Clearance Officer. All requests, emailed or written, should be identified by the title of the information collection.

SUPPLEMENTARY INFORMATION:

Description: This request for an extension is to complete the following data collection activities for MIHOPE-K: (1) A survey with the child's primary caregiver (who will be the mother if she is available), (2) direct assessments of child development, (3) surveys with the child's teacher, (4) a direct assessment of the caregiver, (5) videotaped interactions between the caregiver and child, (6) a caregiver website to provide current contact information, (7) state child welfare records, and (8) school records. In addition to collecting these data, the MIHOPE-K project will continue to maintain up-to-date consent forms for the collection of administrative data. Future information

collection requests and related Federal Register notices will describe future data collection efforts for this project.

Data collected during the kindergarten follow-up study is being used to estimate the effects of MIECHV-funded programs on the following seven domains: (1) Maternal health, (2) child health, (3) child development and school performance, (4) child maltreatment, (5) parenting, (6) crime or domestic violence, and (7) family economic self-sufficiency.

Respondents: The respondents in this extension will include 1.391 families who have not vet participated in the kindergarten follow-up study activities. We have assumed that only 25 percent of respondents will complete the caregiver website. We will also obtain child welfare data from 11 MIHOPE states and school records data from state and local agencies. We have assumed that we will obtain data from 11 states and 5 local education agencies.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents (total over request period)	Number of responses per respondent (total over request period)	Avg. burden per response (in hours)	Total burden (in hours)	Annual burden (in hours)
Burden for previously approved, ongoing data collection					
Survey of caregivers Direct assessments of children Survey of the focal children's teachers Direct assessments of caregivers Videotaped caregiver-child interactions Caregiver website State child welfare records: data file submission School records: data file submission	1,391 1,391 1,391 1,391 2,782 348 11	1 1 1 1 1 1 2 2	0.99 1.33 0.5 0.17 0.25 0.17 15 22.5	1,377 1,850 696 236 696 59 330 720	689 925 348 118 348 30 165 360

Estimated Total Annual Burden Hours: 2,983.

Comments: The Department specifically requests comments on (a) whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Authority: Social Security Act Title V 511 [42 U.S.C. 711]. As extended by the Bipartisan Budget Act of 2018 (Pub. L. 115-123) through FY22.

Mary B. Jones,

ACF/OPRE Certifying Officer.

[FR Doc. 2021-20798 Filed 9-23-21; 8:45 am]

BILLING CODE 4184-74-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [Docket No. FDA-2021-N-0441]

Cardiovascular and Renal 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 Cardiovascular and Renal 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.

DATES: The meeting will be held on December 8, 2021, from 9:30 a.m. to 5 p.m. Eastern Time.

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

Answers to commonly asked questions about FDA advisory committee meetings may be accessed at: https://www.fda.gov/AdvisoryCommittees/AboutAdvisoryCommittees/ucm408555.htm.

FDA is establishing a docket for public comment on this meeting. The docket number is FDA-2021-N-0441. The docket will close on December 7, 2021. Submit either electronic or written comments on this public meeting by December 7, 2021. Please note that late, untimely filed comments will not be considered. Electronic comments must be submitted on or before December 7, 2021. The https:// www.regulations.gov electronic filing system will accept comments until 11:59 p.m. Eastern Time at the end of December 7, 2021. 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.

Comments received on or before November 23, 2021, will be provided to the committee. Comments received after that date will be taken into consideration by FDA. In the event that the meeting is cancelled, FDA will continue to evaluate any relevant applications or information, and consider any comments submitted to the docket, as appropriate.

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): 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—2021—N—0441 for "Cardiovascular and Renal Drugs Advisory Committee; Notice of Meeting; Establishment of a Public Docket; Request for Comments." Received comments, those filed in a timely manner (see ADDRESSES), 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." FDA 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 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 the 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.

FOR FURTHER INFORMATION CONTACT:

Joyce Yu, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2417, Silver Spring, MD 20993-0002, 301-837-7126, Fax: 301-847-8533, email: CRDAC@ fda.hhs.gov, or FDA Advisory Committee Information Line, 1-800-741-8138 (301-443-0572 in the Washington, DC area). A notice in the Federal Register about last minute modifications that impact a previously announced advisory committee meeting cannot always be published quickly enough to provide timely notice. Therefore, you should always check the FDA's website at https://www.fda.gov/ AdvisoryCommittees/default.htm and scroll down to the appropriate advisory committee meeting link, or call the advisory committee information line to learn about possible modifications before coming to the meeting.

SUPPLEMENTARY INFORMATION: Agenda: The meeting presentations will be heard, viewed, captioned, and recorded through an online teleconferencing platform. The committee will discuss new drug application (NDA) 215484, for the Nrf2 activator, bardoxolone methyl capsules, submitted by Reata Pharmaceuticals, Inc. The proposed indication is to slow the progression of chronic kidney disease caused by Alport syndrome in patients 12 years of age and older.

FDA intends to make background material available to the public no later than 2 business days before the meeting. If FDA is unable to post the background material on its website prior to the meeting, the background material will be made publicly available on FDA's

website at the time of the advisory committee meeting. Background material and the link to the online teleconference meeting room will be available at https://www.fda.gov/AdvisoryCommittees/Calendar/default.htm. Scroll down to the appropriate advisory committee meeting link. The meeting will include slide presentations with audio components to allow the presentation of materials in a manner that most closely resembles an in-person advisory committee meeting.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. All electronic and written submissions submitted to the Docket (see ADDRESSES) on or before November 23, 2021, will be provided to the committee. Oral presentations from the public will be scheduled between approximately 2:30 p.m. and 3:30 p.m. Eastern Time. Those individuals interested in making formal oral presentations should notify the contact person and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation on or before November 15, 2021. Time allotted for each presentation may be limited. If the number of registrants requesting to speak is greater than can be reasonably accommodated during the scheduled open public hearing session, FDA may conduct a lottery to determine the speakers for the scheduled open public hearing session. The contact person will notify interested persons regarding their request to speak by November 16, 2021.

For press inquiries, please contact the Office of Media Affairs at *fdaoma@ fda.hhs.gov* or 301–796–4540.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with disabilities. If you require accommodations due to a disability, please contact Joyce Yu (see FOR FURTHER INFORMATION CONTACT) at least 7 days in advance of the meeting.

FDA is committed to the orderly conduct of its advisory committee meetings. Please visit our website at https://www.fda.gov/Advisory Committees/AboutAdvisoryCommittees/ucm111462.htm for procedures on public conduct during advisory committee meetings.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2). Dated: September 17, 2021.

Lauren K. Roth,

 $Acting \ Principal \ Associate \ Commissioner for \ Policy.$

[FR Doc. 2021–20733 Filed 9–23–21; 8:45 am]

BILLING CODE 4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2021-N-0973]

Revocation of Three Authorizations of Emergency Use of In Vitro Diagnostic Devices for Detection and/or Diagnosis of COVID-19; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the revocation of the Emergency Use Authorizations (EUAs) (the Authorizations) issued to Gravity Diagnostics, LLC (Gravity) for the Gravity Diagnostics COVID-19 Assay, Materials and Machines Corporation of America (DBA MatmaCorp, Inc.) (Matmacorp) for the MatMaCorp COVID-19 2SF Test, and Guardant Health Inc. (Guardant) for the Guardant-19. FDA revoked Gravity's Authorization on July 21, 2021, Matmacorp's Authorization on August 3, 2021, and Guardant's Authorization on August 6, 2021, under the Federal Food, Drug, and Cosmetic Act (FD&C Act). The revocations, which include an explanation of the reasons for each revocation, are reprinted in this document.

DATES: Gravity's Authorization is revoked as of July 21, 2021. Matmacorp's Authorization is revoked as of August 3, 2021. Guardant's Authorization is revoked as of August 6, 2021.

ADDRESSES: Submit written requests for single copies of the revocations to the Office of Counterterrorism and Emerging Threats, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 1, Rm. 4338, Silver Spring, MD 20993–0002. Send one self-addressed adhesive label to assist that office in processing your request or include a Fax number to which the revocations may be sent. See the SUPPLEMENTARY INFORMATION section for electronic access to the revocations.

FOR FURTHER INFORMATION CONTACT: Jennifer J. Ross, Office of

Counterterrorism and Emerging Threats, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 1, Rm. 4332, Silver Spring, MD 20993–0002, 240–402–8155 (this is not a toll-free number).

SUPPLEMENTARY INFORMATION:

I. Background

Section 564 of the FD&C Act (21 U.S.C. 360bbb-3) allows FDA to strengthen the public health protections against biological, chemical, nuclear, and radiological agents. Among other things, section 564 of the FD&C Act allows FDA to authorize the use of an unapproved medical product or an unapproved use of an approved medical product in certain situations. On June 1, 2020, FDA issued an EUA to Gravity for the Gravity Diagnostics COVID-19 Assay. Notice of the issuance of this Authorization was published in the Federal Register on November 20, 2020 (85 FR 74346), as required by section 564(h)(1) of the FD&C Act. On August 21, 2020, FDA issued an EUA to Guardant for the Guardant-19. Notice of the issuance of this Authorization was published in the Federal Register on November 20, 2020 (85 FR 74346), as required by section 564(h)(1) of the FD&C Act. On December 17, 2020, FDA issued an EUA to Matmacorp, for the MatMaCorp COVID-19 2SF Test. Notice of the issuance of this Authorization was published in the Federal Register on April 23, 2021 (86 FR 21749), as required by section 564(h)(1) of the FD&C Act. The authorization of a device for emergency use under section 564 of the FD&C Act may, pursuant to section 564(g)(2) of the FD&C Act, be revoked when the criteria under section 564(c) of the FD&C Act for issuance of such authorization are no longer met (section 564(g)(2)(B) of the FD&C Act), or other circumstances make such revocation appropriate to protect the public health or safety (section 564(g)(2)(C) of the FD&C Act).

II. EUA Revocation Requests

On March 11, 2021, and reconfirmed July 12, 2021, Gravity requested the revocation of, and on July 21, 2021, FDA revoked, the Authorization for the Gravity Diagnostics COVID-19 Assay. Because Gravity notified FDA that it is no longer using the Gravity Diagnostics COVID-19 Assay and requested FDA revoke the Authorization, FDA has determined that it is appropriate to protect the public health or safety to revoke this Authorization. On July 29, 2021, Matmacorp requested the revocation of, and on August 3, 2021, FDA revoked, the Authorization for the MatMaCorp COVID-19 2SF Test. Because Matmacorp notified FDA that it