

I. General Information

A. Does this action apply to me?

You may be potentially affected by this action if you are an agricultural producer, food manufacturer, or pesticide manufacturer. The following list of North American Industrial Classification System (NAICS) codes is not intended to be exhaustive, but rather provides a guide to help readers determine whether this document applies to them. Potentially affected entities may include:

- Crop production (NAICS code 111).
- Animal production (NAICS code 112).
- Food manufacturing (NAICS code 311).

B. What should I consider as I prepare my comments for EPA?

1. *Submitting CBI.* Do not submit this information to EPA through [regulations.gov](https://www.epa.gov/regulations.gov) or email. Clearly mark the part or all of the information that you claim to be CBI. For CBI information in a disk or CD-ROM that you mail to EPA, mark the outside of the disk or CD-ROM as CBI and then identify electronically within the disk or CD-ROM the specific information that is claimed as CBI. In addition to one complete version of the comment that includes information claimed as CBI, a copy of the comment that does not contain the information claimed as CBI must be submitted for inclusion in the public docket. Information so marked will not be disclosed except in accordance with procedures set forth in 40 CFR part 2.

2. *Tips for preparing your comments.* When preparing and submitting your comments, see the commenting tips at <https://www.epa.gov/dockets/commenting-epa-dockets>.

II. Registration Applications

EPA has received applications to register pesticide products containing active ingredients not included in any currently registered pesticide products. Pursuant to the provisions of FIFRA section 3(c)(4) (7 U.S.C. 136a(c)(4)), EPA is hereby providing notice of receipt and opportunity to comment on these applications. Notice of receipt of these applications does not imply a decision by the Agency on these applications. For actions being evaluated under EPA's public participation process for registration actions, there will be an additional opportunity for public comment on the proposed decisions. Please see EPA's public participation website for additional information on this process ([http://www2.epa.gov/pesticide-registration/public-](http://www2.epa.gov/pesticide-registration/public-participation-process-registration-actions)

[participation-process-registration-actions](http://www2.epa.gov/pesticide-registration/public-participation-process-registration-actions)).

A. Notice of Receipt—New Active Ingredients

1. *File Symbols:* 52991–GL and 51934–ET. *Docket ID number:* EPA–HQ–OPP–2021–0651. *Applicant:* Spring Regulatory Sciences on behalf of Bedoukian Research, Inc., 21 Finance Drive, Danbury, CT 06810–4192. *Product name:* Bedoukian Serricornin Technical Pheromone and Cidetrak CB. *Active ingredient:* Mating disruptant, rac-4S,6S,7S-serricornin at 63.0% and 5.0% by weight, respectively. *Proposed classification/Use:* Mating disruption for cigarette beetle, lasioderma serricorne. *Contact:* BPPD.

2. *File Symbol:* 91283–RE. *Docket ID number:* EPA–HQ–OPP–2021–0786. *Applicant:* Amoéba SA 38 Avenue des Frères Montgolfier, F–69680 Chassieu, France (c/o SciReg, Inc. 12733 Director's Loop, Woodbridge, VA 22192). *Product name:* Amoéba EP #2. *Active ingredient:* Antimicrobial—*Willaertia Magna* C2c.Maky at 1%. *Proposed use:* For control of microbial slime (bioslime), microbially induced non-public health corrosion and non-public health general microbial flora in cooling towers waters. *Contact:* BPPD.

3. *File Symbol:* 91283–RG. *Docket ID number:* EPA–HQ–OPP–2021–0786. *Applicant:* Amoéba SA 38 Avenue des Frères Montgolfier, F–69680 Chassieu, France (c/o SciReg, Inc. 12733 Director's Loop, Woodbridge, VA 22192). *Product name:* Amoéba EP #1. *Active ingredient:* Antimicrobial—*Willaertia Magna* C2c.Maky at 1%. *Proposed use:* For control of microbial slime (bioslime), microbially induced non-public health corrosion and non-public health general microbial flora in cooling towers waters. *Contact:* BPPD.

4. *File Symbol:* 91283–RN. *Docket ID number:* EPA–HQ–OPP–2021–0786. *Applicant:* Amoéba SA 38 Avenue des Frères Montgolfier, F–69680 Chassieu, France (c/o SciReg, Inc. 12733 Director's Loop, Woodbridge, VA 22192). *Product name:* Amoéba TGAI. *Active ingredient:* Antimicrobial—*Willaertia Magna* C2c.Maky at 1%. *Proposed use:* For control of microbial slime (bioslime), microbially induced non-public health corrosion and non-public health general microbial flora in cooling towers waters. *Contact:* BPPD.

5. *File Symbol:* 91283–RR. *Docket ID number:* EPA–HQ–OPP–2021–0786. *Applicant:* Amoéba SA 38 Avenue des Frères Montgolfier, F–69680 Chassieu, France (c/o SciReg, Inc. 12733 Director's Loop, Woodbridge, VA 22192). *Product name:* Amoéba EP #3. *Active ingredient:* Antimicrobial—*Willaertia Magna*

C2c.Maky at 1%. *Proposed use:* For control of microbial slime (bioslime), microbially induced non-public health corrosion and non-public health general microbial flora in cooling towers waters. *Contact:* BPPD.

Authority: 7 U.S.C. 136 *et seq.*

Dated: December 7, 2021.

Delores Barber,

Director, Information Technology and Resources Management Division, Office of Program Support.

[FR Doc. 2021–27301 Filed 12–16–21; 8:45 am]

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ENVIRONMENTAL PROTECTION AGENCY

[ER–FRL–9059–8]

Environmental Impact Statements; Notice of Availability

Responsible Agency: Office of Federal Activities, General Information 202–564–5632 or <https://www.epa.gov/nepa>.

Weekly receipt of Environmental Impact Statements (EIS)

Filed December 6, 2021 10 a.m. EST

Through December 13, 2021 10 a.m. EST

Pursuant to 40 CFR 1506.9.

Notice

Section 309(a) of the Clean Air Act requires that EPA make public its comments on EISs issued by other Federal agencies. EPA's comment letters on EISs are available at: <https://cdxnodengn.epa.gov/cdx-enepa-public/action/eis/search>.

EIS No. 20210184, Final, USFS, OR, Stella Landscape Restoration Project, Review Period Ends: 01/31/2022, Contact: Michelle Calvert 541–441–7059.

Dated: December 13, 2021.

Cindy S. Barger,

Director, NEPA Compliance Division, Office of Federal Activities.

[FR Doc. 2021–27336 Filed 12–16–21; 8:45 am]

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ENVIRONMENTAL PROTECTION AGENCY

[FRL–9328–01–ORD]

Human Studies Review Board (HSRB); Notification of Public Meetings

AGENCY: Environmental Protection Agency (EPA).

ACTION: Notice of public meeting.

SUMMARY: The Environmental Protection Agency (EPA), Office of Research and Development (ORD), gives notice of the

2022 public meetings of the Human Studies Review Board (HSRB). The HSRB provides advice, information, and recommendations on issues related to scientific and ethical aspects of third-party human subjects' research that are submitted to the Office of Pesticide Programs (OPP) to be used for regulatory purposes.

DATES: Four three-day virtual public meetings will be held on:

1. January 25–27, 2022;
2. April 26–28, 2022;
3. July 19–21, 2022; and
4. October 25–27, 2022.

Meetings will be held each day from 1 p.m. to 5 p.m. Eastern Time. For each meeting, separate subsequent follow-up meetings are planned for the HSRB to finalize reports from the three-day meetings. These meetings will be held from 2 p.m. to 4 p.m. Eastern time on the following dates: March 17, 2022; June 16, 2022; September 14, 2022; and December 14, 2022.

ADDRESSES: These meetings are open to the public and will be conducted entirely virtually and by telephone. For detailed access information and meeting materials please visit the HSRB website: <https://www.epa.gov/osa/human-studies-review-board>.

FOR FURTHER INFORMATION CONTACT: Any member of the public who wished to receive further information should contact the HSRB Designated Federal Official (DFO), Tom Tracy, via phone/voicemail at: 919–541–4334; or via email at: tracy.tom@epa.gov.

SUPPLEMENTARY INFORMATION:

Background

The HSRB is a Federal advisory committee operating in accordance with the Federal Advisory Committee Act 5 U.S.C. App.2 section 9. The HSRB provides advice, information, and recommendations on issues related to scientific and ethical aspects of third-party human subjects research that are submitted to the Office of Pesticide Programs (OPP) to be used for regulatory purposes.

Meeting access: These meetings will be open to the public. The full agenda with access information and meeting materials will be available seven calendar days prior to the start of each meeting at the HSRB website: <https://www.epa.gov/osa/human-studies-review-board>. For questions on document availability, or if you do not have access to the internet, consult with the DFO, Tom Tracy listed under **FOR FURTHER INFORMATION CONTACT**.

Special Accommodations. For information on access or services for individuals with disabilities, or to

request accommodation of a disability, please contact the DFO listed under **FOR FURTHER INFORMATION CONTACT** at least 10 days prior to each meeting to give EPA as much time as possible to process your request.

How may I participate in this meeting?

The HSRB encourages the public's input. You may participate in these meetings by following the instructions in this section.

1. *Oral comments.* To pre-register to make oral comments, please contact the DFO, Tom Tracy, listed under **FOR FURTHER INFORMATION CONTACT**. Requests to present oral comments during the meetings will be accepted up to Noon Eastern Time, seven calendar days prior to each meeting date. To the extent that time permits, interested persons who have not pre-registered may be permitted by the HSRB Chair to present oral comments during the meetings at the designated time on the agenda. Oral comments before the HSRB are generally limited to five minutes per individual or organization. If additional time is available, further public comments may be possible.

2. *Written comments.* For the Board to have the best opportunity to review and consider your comments as it deliberates, you should submit your comments prior to the meetings via email by Noon Eastern Time, seven calendar days prior to each meeting date. If you submit comments after these dates, those comments will be provided to the HSRB members, but you should recognize that the HSRB members may not have adequate time to consider your comments prior to their discussion. You should submit your comments to the DFO, Tom Tracy listed under **FOR FURTHER INFORMATION CONTACT**. There is no limit on the length of written comments for consideration by the HSRB.

Topics for discussion. The agenda and meeting materials will be available seven calendar days in advance of each meeting at <https://www.epa.gov/osa/human-studies-review-board>.

Meeting minutes and final reports. Minutes of these meetings, summarizing the topics discussed and recommendations made by the HSRB, will be released within 90 calendar days of each meeting. These minutes will be available at <https://www.epa.gov/osa/human-studies-review-board>. In addition, information regarding the HSRB's Final Reports, will be found at <https://www.epa.gov/osa/human-studies-review-board> or can be

requested from Tom Tracy listed under **FOR FURTHER INFORMATION CONTACT**.

Mary Ross,

Director, Office of Science Advisor, Policy and Engagement.

[FR Doc. 2021–27396 Filed 12–16–21; 8:45 am]

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FEDERAL DEPOSIT INSURANCE CORPORATION

Designated Reserve Ratio for 2022

AGENCY: Federal Deposit Insurance Corporation.

ACTION: Notice of Designated Reserve Ratio for 2022.

SUMMARY: Pursuant to the Federal Deposit Insurance Act (FDI Act), the Board of Directors of the Federal Deposit Insurance Corporation designates that the Designated Reserve Ratio (DRR) for the Deposit Insurance Fund shall remain at 2 percent for 2022. The Board is publishing this notice as required by section 7(b)(3)(A)(i) the FDI Act (12 U.S.C. 1817(b)(3)(A)(i)).

FOR FURTHER INFORMATION CONTACT: Ashley Mihalik, Chief, Banking and Regulatory Policy Section, Division of Insurance and Research, (202) 898–3793, amihalik@fdic.gov; Daniel Hoople, Acting Chief, Fund Analysis and Pricing Section, Division of Insurance and Research, (202) 898–3835, dhoople@fdic.gov; or Nefretete Smith, Counsel, Legal Division, (202) 898–6851, nefsmith@fdic.gov.

SUPPLEMENTARY INFORMATION: Pursuant to the FDI Act, the Board designates that the DRR for the Deposit Insurance Fund shall remain at 2 percent for 2022. The Board is publishing this notice as required by section 7(b)(3)(A)(i) the FDI Act (12 U.S.C. 1817(b)(3)(A)(i)). There is no need to amend 12 CFR 327.4(g), the section of the FDIC's regulations which sets forth the DRR, because the DRR for 2022 is the same as the current DRR.

Dated at Washington, DC, on December 14, 2021.

Federal Deposit Insurance Corporation.

By order of the Board of Directors.

James P. Sheesley,

Assistant Executive Secretary.

[FR Doc. 2021–27382 Filed 12–16–21; 8:45 am]

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