mechanisms to audit access attempts and access granted into these areas.

#### RECORD ACCESS PROCEDURES:

All requests for access to records must be in writing and should be submitted to the OASCR FOIA Officer, 1400 Independence Avenue SW, Washington, DC 20250; or by email at USDAFOIA@ usda.gov. In accordance with 7 CFR part 1, subpart G, § 1.112 (Procedures for requests pertaining to individual records in a record system), the request must include the full name of the individual making the request; the name of the system of records; and preference of inspection, in person or by mail. In accordance with 7 CFR 1.113, prior to inspection of the records, the requester shall present sufficient identification (e.g., driver's license, employee identification card, social security card, credit cards) to establish that the requester is the individual to whom the records pertain. In addition, if an individual submitting a request for access wishes to be supplied with copies of the records by mail, the requester must include with his or her request sufficient data for the agency to verify the requester's identity.

### CONTESTING RECORD PROCEDURES:

Individuals seeking to contest or amend records maintained in this system of records must direct their request to the address indicated in the "RECORD ACCESS PROCEDURES" paragraph, above and must follow the procedures set forth in 7 CFR part 1, subpart G, § 1.116 (Request for correction or amendment to record). All requests must state clearly and concisely what record is being contested, the reasons for contesting it, and the proposed amendment to the record.

#### **NOTIFICATION PROCEDURES:**

Individuals may be notified if a record in this system of records pertains to them when the individuals request information utilizing the same procedures as those identified in the "RECORD ACCESS PROCEDURES" paragraph, above.

#### **EXEMPTIONS PROMULGATED FOR THE SYSTEM:**

None.

#### HISTORY:

None.

[FR Doc. 2021–17569 Filed 8–16–21; 8:45 am] BILLING CODE 3410–9R–P

#### **DEPARTMENT OF AGRICULTURE**

Animal and Plant Health Inspection Service

[Docket No. APHIS-2020-0023]

BASF Corporation; Availability of a Draft Plant Pest Risk Assessment and Draft Environmental Assessment for Determination of Nonregulated Status of Plant-Parasitic Nematode-Protected and Herbicide Tolerant Soybean

**AGENCY:** Animal and Plant Health Inspection Service, USDA.

**ACTION:** Notice.

**SUMMARY:** We are advising the public that the Animal and Plant Health Inspection Service has prepared a draft plant pest risk assessment and draft environmental assessment regarding a request from BASF Corporation seeking a determination of nonregulated status for soybean event GMB151, which has been developed using genetic engineering for resistance to the plantparasitic nematode, soybean cyst nematode (Heterodera glycines), and for tolerance to 4-hydroxyphenylpyruvate dioxygenase (HPPD-4) inhibitor herbicides. We are making these documents available for public review and comment.

**DATES:** We will consider all comments that we receive on or before September 16, 2021

**ADDRESSES:** You may submit comments by either of the following methods:

- Federal eRulemaking Portal: Go to www.regulations.gov. Enter APHIS—2020—0023 in the Search field. Select the Documents tab, then select the Comment button in the list of documents.
- Postal Mail/Commercial Delivery: Send your comment to Docket No. APHIS-2020-0023, Regulatory Analysis and Development, PPD, APHIS, Station 3A-03.8, 4700 River Road, Unit 118, Riverdale, MD 20737-1238.

The petition, draft environmental assessment, draft plant pest risk assessment, and any comments we receive on this docket may be viewed at www.regulations.gov, or in our reading room, which is located in Room 1620 of the USDA South Building, 14th Street and Independence Avenue SW, Washington, DC. Normal reading room hours are 8 a.m. to 4:30 p.m., Monday through Friday, except holidays. To be sure someone is there to help you, please call (202) 7997039 before coming.

Supporting documents for this petition are also available on the APHIS website at https://www.aphis.usda.gov/aphis/ourfocus/biotechnology/permits-

notifications-petitions/petitions/petition-status.

FOR FURTHER INFORMATION CONTACT: Ms. Cindy Eck, Biotechnology Regulatory Services, APHIS, 4700 River Road, Unit 147, Riverdale, MD 20737–1236; (301) 851–3892, email: cynthia.a.eck@usda.gov.

**SUPPLEMENTARY INFORMATION:** Under the authority of the plant pest provisions of the Plant Protection Act (7 U.S.C. 7701 et seq.), the regulations in 7 CFR part 340, "Movement of Organisms Modified or Produced Through Genetic Engineering," regulate, among other things, the importation, interstate movement, or release into the environment of organisms modified or produced through genetic engineering that are plant pests or pose a plausible

plant pest risk.

The petition for nonregulated status described in this notice is being evaluated under the version of the regulations effective at the time that it was received. The Animal and Plant Health Inspection Service (APHIS) issued a final rule, published in the Federal Register on May 18, 2020 (85 FR 29790-29838, Docket No. APHIS-2018–0034),1 revising 7 CFR part 340; however, the final rule is being implemented in phases. The new Regulatory Status Review (RSR) process, which replaces the petition for determination of nonregulated status process, became effective on April 5, 2021 for corn, soybean, cotton, potato, tomato, and alfalfa. The RSR process is effective for all crops as of October 1, 2021. However, "[u]ntil RSR is available for a particular crop . . . APHIS will continue to receive petitions for determination of nonregulated status for the crop in accordance with the [legacy] regulations at 7 CFR 340.6." (85 FR 29815). This petition for a determination of nonregulated status is being evaluated in accordance with the regulations at 7 CFR 340.6 (2020) as it was received by APHIS on November

BASF Corporation (BASF) has submitted a petition (APHIS Petition Number 19–317–01p) to APHIS seeking a determination of nonregulated status under 7 CFR part 340, for soybean event GMB151 which has been developed using genetic engineering for resistance to the plant-parasitic nematode, soybean cyst nematode (*Heterodera glycines*), and for tolerance to 4-hydroxyphenylpyruvate dioxygenase (HPPD-4) inhibitor herbicides. The petition states that GMB151 soybean is

 $<sup>^{\</sup>rm 1}\,\rm To$  view the final rule, go to www.regulations.gov and enter APHIS–2018–0034 in the Search field.

unlikely to pose a plant pest risk and, therefore, should not be regulated under APHIS' regulations in 7 CFR part 340.

According to our process <sup>2</sup> for soliciting public comment when considering petitions for determination of nonregulated status of organisms developed using genetic engineering, APHIS accepts written comments regarding a petition once APHIS deems the petition complete. On May 28, 2020, APHIS announced in the **Federal Register** <sup>3</sup> (85 FR 32004–32005, Docket No. APHIS–2020–0023) the availability of the BASF petition for public comment. APHIS solicited comments on the petition for 60 days ending July 27, 2020.

APHIS received nine comments during the comment period. They were from the agricultural and private sectors. Five comments generally supported BASF's petition, while four expressed objections to crops developed or modified through genetic engineering.

After public comments are received on a completed petition, APHIS evaluates those comments and then provides a second opportunity for public involvement in our decision-making process. According to our public review process (see footnote 2), the second opportunity for public involvement follows one of two approaches, as described below.

If APHIS decides, based on its review of the petition and its evaluation and analysis of comments received during the 60-day public comment period on the petition, that the petition involves an organism that raises no substantive new issues, APHIS will follow Approach 1 for public involvement. Under Approach 1, APHIS prepares and announces in the Federal Register the availability of APHIS' preliminary regulatory determination along with its draft EA, preliminary finding of no significant impact (FONSI), and its draft plant pest risk assessment (PPRA) for a 30-day public review period. APHIS will evaluate any information received related to the petition and its supporting documents during the 30-day public review period. If APHIS determines that no substantive information has been

received that would warrant APHIS altering its preliminary regulatory determination or FONSI, or substantially change the analysis of impacts in the EA, our preliminary regulatory determination will become final and effective upon notification of the public through an announcement on our website. No further **Federal Register** notice will be published announcing the final regulatory determination.

Under Approach 2, if APHIS decides, based on its review of the petition and its evaluation and analysis of comments received during the 60-day public comment period on the petition, that the petition involves an organism that raises substantive new issues, APHIS first solicits written comments from the public on a draft EA and draft PPRA for a 30-day comment period through the publication of a Federal Register notice. Then, after reviewing and evaluating the comments on the draft EA and draft PPRA and other information, APHIS will revise the draft PPRA as necessary. It will then prepare a final EA, and based on the final EA, a National Environmental Policy Act (NEPA) decision document (either a FONSI or a notice of intent to prepare an environmental impact statement).

For this petition, we will be following Approach 2.

As part of our decision-making process regarding an organism's regulatory status, APHIS prepared a PPRA to assess the plant pest risk of the organism, and an EA to evaluate potential impacts on the human environment. This will provide the Agency and the public with a review and analysis of any potential environmental impacts that may result if the petition request is approved.

APHIS' draft PPRA compared the pest risk posed by soybean event GMB151 with that of the unmodified variety from which it was derived. The draft PPRA concluded that soybean event GMB151 is unlikely to pose an increased plant pest risk compared to the unmodified soybean.

The draft EA evaluated potential impacts that may result from the commercial production of GMB151 soybean, to include potential impacts on conventional and organic soybean production; the acreage and area required for U.S. soybean production; agronomic practices and inputs; the physical environment; biological resources; human health and worker safety; animal health and welfare; and socioeconomic impacts. No significant impacts were identified with the production and marketing of GMB151 soybean.

The draft EA was prepared in accordance with (1) NEPA, as amended (42 U.S.C. 4321 et seq.), (2) regulations of the Council on Environmental Quality for implementing the procedural provisions of NEPA (40 CFR parts 1500–1508), (3) USDA regulations implementing NEPA (7 CFR part 1b), and (4) APHIS' NEPA Implementing Procedures (7 CFR part 372).

We are making available for a 30-day review period our draft EA and draft PPRA. These documents are available as indicated under ADDRESSES and FOR FURTHER INFORMATION CONTACT above. Copies of these documents may also be obtained from the person listed under FOR FURTHER INFORMATION CONTACT.

After the 30-day review period closes, APHIS will review and evaluate any information received during the 30-day review period.

Authority: 7 U.S.C. 7701–7772 and 7781–7786; 31 U.S.C. 9701; 7 CFR 2.22, 2.80, and 371.3.

Done in Washington, DC, this 11th day of August 2021.

#### Michael Watson,

Acting Administrator, Animal and Plant Health Inspection Service.

[FR Doc. 2021–17558 Filed 8–16–21; 8:45 am]

BILLING CODE 3410-34-P

## **COMMISSION ON CIVIL RIGHTS**

# Notice of Public Meetings of the Minnesota Advisory Committee

**AGENCY:** U.S. Commission on Civil Rights.

**ACTION:** Announcement of meeting.

SUMMARY: Notice is hereby given, pursuant to the provisions of the rules and regulations of the U.S. Commission on Civil Rights (Commission) and the Federal Advisory Committee Act that the Minnesota Advisory Committee (Committee) will hold a meeting via the online platform WebEx on Tuesday, August 24, 2021 at 12:00 p.m. Central Time. The purpose of the meeting is to discuss a memorandum on civil rights concerns in the state.

**DATES:** The meeting will be held on:

• Tuesday, August 24, 2021, at 12:00 p.m. Central Time

Web link: https://civilrights.webex.com/ civilrights/j.php?MTID= m16213078bd3f943a55c68fe 7491c75ad

Join by phone: 800–360–9505 USA Toll Free

Access code: 199 660 9075

#### FOR FURTHER INFORMATION CONTACT:

David Barreras, Designated Federal Officer, at *dbarreras@usccr.gov* or (202) 656–8937.

<sup>&</sup>lt;sup>2</sup> On March 6, 2012, APHIS published in the Federal Register (77 FR 13258–13260, Docket No. APHIS–2011–0129) a notice describing our public review process for soliciting public comments and information when considering petitions for determinations of nonregulated status for organisms developed using genetic engineering. To view the notice, go to <a href="https://www.regulations.gov">www.regulations.gov</a> and enter APHIS–2011–0129 in the Search field.

<sup>&</sup>lt;sup>3</sup> To view the notice, its supporting documents, and the comments that we received, go to www.regulations.gov and enter APHIS-2020-0023 in the Search field.