

IV. Participating in the Public Hearing

Registration: To register to attend the virtual public hearing, on “Scientific Data and Information Related to the Residue of Carcinogenic Concern for the New Animal Drug Carbadox; Public Hearing; Request for Comments” please register at <https://fda.zoomgov.com/j/1600135012?pwd=MFdjMW9FRXg4RGllc3FHVVhkWVAYzZ09> by March 9, 2022. If you have any questions, you can contact CarbadoxPublicHearing2022@fda.hhs.gov (See **DATES** and **ADDRESSES**). Please provide complete contact information for each attendee, including name, title, affiliation, address, email, and telephone.

Request for Oral Presentations: During online registration, you may indicate if you wish to make a formal presentation (with accompanying slide deck) or present oral comments during the public hearing session (with no slide deck). If you decide you wish to make a presentation after registering online, you may submit a request to CarbadoxPublicHearing2022@fda.hhs.gov. All requests to make presentations must be received by February 18, 2022. FDA will do its best to accommodate requests to make public presentations. Individuals and organizations with common interests are urged to consolidate or coordinate their presentations. FDA will determine the amount of time allotted to each presenter and the approximate time each presentation is to begin and will select and notify participants by February 23, 2022.

If selected for a formal oral presentation (with a slide deck), each presenter must submit an electronic copy of their presentation (PowerPoint or PDF) to CarbadoxPublicHearing2022@fda.hhs.gov with the subject line “Scientific Data and Information Related to the Residue of Carcinogenic Concern for the New Animal Drug Carbadox; Public Hearing; Request for Comments” on or before March 4, 2022. No commercial or promotional material will be permitted to be presented or distributed at the public hearing.

Persons notified that they will be presenters are encouraged to be online early. Actual presentation times may vary based on how the hearing progresses in real time. An agenda for the hearing and any other background materials will be made available no later than 5 days before the hearing at <https://www.fda.gov/animal-veterinary/workshops-conferences-meetings/part-15-public-hearing-scientific-data-and->

information-related-residue-carcinogenic-concern-new.

Transcripts: Please be advised that as soon as a transcript of the public hearing is available, it will be accessible at <https://www.regulations.gov>. It may also be viewed at the Dockets Management Staff (see **ADDRESSES**). A link to the transcript will also be available on the Agency’s website at <https://www.fda.gov/animal-veterinary/workshops-conferences-meetings/part-15-public-hearing-scientific-data-and-information-related-residue-carcinogenic-concern-new>.

Dated: January 7, 2022.

Lauren K. Roth,

Associate Commissioner for Policy.

[FR Doc. 2022–00475 Filed 1–12–22; 8:45 am]

BILLING CODE 4164–01–P

DEPARTMENT OF THE INTERIOR

National Indian Gaming Commission

25 CFR Part 559

RIN 3141–AA76

Facility License; Correction

AGENCY: National Indian Gaming Commission, Department of the Interior.

ACTION: Proposed rule; correction.

SUMMARY: This document corrects the preamble to a proposed rule published in the **Federal Register** of December 1, 2021, regarding Facility Licenses. The document contained incorrect dates for submitting comments. This correction clarifies that comments are due January 31, 2022.

FOR FURTHER INFORMATION CONTACT: Michael Hoenig, 202–632–7003.

SUPPLEMENTARY INFORMATION:

Correction

In the **Federal Register** of December 1, 2021, in FR Doc. 2021–25845, on page 68200, in the third column, change the **DATES** caption to read:

DATES: Written comments on this proposed rule must be received on or before January 31, 2022.

Dated: January 6, 2022.

Michael Hoenig,

General Counsel.

[FR Doc. 2022–00625 Filed 1–12–22; 8:45 am]

BILLING CODE 7565–01–P

ENVIRONMENTAL PROTECTION AGENCY

40 CFR Part 52

[EPA–R04–OAR–2021–0428; FRL–9374–01–R4]

Finding of Failure To Attain the 2010 Sulfur Dioxide Standard; Tennessee; Sullivan County Nonattainment Area

AGENCY: Environmental Protection Agency (EPA).

ACTION: Proposed rule.

SUMMARY: The Environmental Protection Agency (EPA) is proposing to determine that the Sullivan County, Tennessee sulfur dioxide (SO₂) nonattainment area failed to attain the 2010 1-hour SO₂ primary National Ambient Air Quality Standard (NAAQS or standard) by the applicable attainment date of October 4, 2018, based upon a weight of evidence analysis of available quality-assured and certified SO₂ ambient air monitoring data and SO₂ emissions data from January 2015 through December 2017. If EPA finalizes this determination as proposed, the State of Tennessee will be required to submit revisions to the Tennessee State Implementation Plan (SIP) that, among other elements, provide for expeditious attainment of the 2010 SO₂ standard.

DATES: Comments must be received on or before February 14, 2022.

ADDRESSES: Submit your comments, identified by Docket ID No. EPA–R04–OAR–2021–0428 at <https://www.regulations.gov>. Follow the online instructions for submitting comments. Once submitted, comments cannot be edited or removed from *Regulations.gov*. EPA may publish any comment received to its public docket. Do not submit electronically any information you consider to be Confidential Business Information (CBI) or other information whose disclosure is restricted by statute. Multimedia submissions (audio, video, etc.) must be accompanied by a written comment. The written comment is considered the official comment and should include discussion of all points you wish to make. EPA will generally not consider comments or comment contents located outside of the primary submission (*i.e.*, on the web, cloud, or other file sharing system). For additional submission methods, the full EPA public comment policy, information about CBI or multimedia submissions, and general guidance on making effective comments, please visit <https://www2.epa.gov/dockets/commenting-epa-dockets>.

FOR FURTHER INFORMATION CONTACT: Evan Adams, Air Regulatory