

for public comment on any proposed risk management actions.

For more information about the TSCA risk evaluation process for existing chemicals, go to <https://www.epa.gov/assessing-and-managing-chemicals-under-tsca>.

(Authority: 15 U.S.C. 2601 *et seq.*)

**Douglas M. Troutman,**  
Assistant Administrator, Office of Chemical Safety and Pollution Prevention.

[FR Doc. 2026-16263 Filed 8-7-26; 8:45 am]

**BILLING CODE 6560-50-P**

## EQUAL EMPLOYMENT OPPORTUNITY COMMISSION

### Commission Meeting—Sunshine Act Notice

**TIME AND DATE:** Tuesday, August 11, 2026, 1:00 p.m. Eastern Time or earlier.

**PLACE:** The meeting will be held in-person.

**STATUS:** The meeting will be closed to the public.

#### **MATTERS TO BE CONSIDERED:**

The following items will be considered at the meeting:

- Pending Litigation Recommendation(s)

**Note:** The Legal Counsel has certified that, in the Legal Counsel's opinion, the Commission meeting scheduled for August 11, 2026, concerning a recommendation to participate in civil litigation may properly be closed under the 3rd, 6th, 7th, and 10th exemptions to the Government in the Sunshine Act, 5 U.S.C. 552b(10), and Commission regulations at 29 CFR 1612.13(a).

Please telephone (202) 921-2750, or email [commissionmeetingcomments@eeoc.gov](mailto:commissionmeetingcomments@eeoc.gov) at any time for information on this meeting.

**CONTACT PERSON FOR MORE INFORMATION:** Raymond Windmiller, Executive Officer, (202) 921-2705.

Dated: August 5, 2026.

**Raymond D. Windmiller,**  
Executive Officer, Executive Secretariat.

[FR Doc. 2026-16244 Filed 8-6-26; 4:15 pm]

**BILLING CODE 6570-01-P**

## FARM CREDIT ADMINISTRATION

### Sunshine Act Meetings

**TIME AND DATE:** 10 a.m., Thursday, August 13, 2026.

**PLACE:** You may observe this meeting in person at 1501 Farm Credit Drive, McLean, Virginia 22102-5090, or virtually. If you would like to observe,

at least 24 hours in advance, visit [FCA.gov](https://www.fca.gov), select "Newsroom," then select "Events." From there, access the linked "Instructions for board meeting visitors" and complete the described registration process.

**STATUS:** This meeting will be open to the public.

**MATTERS TO BE CONSIDERED:** The following matters will be considered:

- Annual Report on the Farm Credit System's Young, Beginning, and Small Farmers and Ranchers Mission Performance
- Office of Secondary Market Oversight Periodic Report

#### **CONTACT PERSON FOR MORE INFORMATION:**

If you need more information or assistance for accessibility reasons, or have questions, contact Ashley Waldron, Secretary to the Board. Telephone: 703-883-4009. TTY: 703-883-4056.

**Ashley Waldron,**  
Secretary to the Board.

[FR Doc. 2026-16200 Filed 8-6-26; 11:15 am]

**BILLING CODE 6705-01-P**

## FEDERAL COMMUNICATIONS COMMISSION

[FR ID: 361207]

### Privacy Act of 1974; Matching Program

**AGENCY:** Federal Communications Commission.

**ACTION:** Notice of a new matching program.

**SUMMARY:** In accordance with the Privacy Act of 1974, as amended ("Privacy Act"), this document announces a new computer matching program the Federal Communications Commission (FCC or Commission or Agency) and the Universal Service Administrative Company (USAC) will conduct with the Iowa Department of Health and Human Services. The purpose of this matching program is to verify the eligibility of applicants to and subscribers of Lifeline, which is administered by USAC under the direction of the FCC. More information about this program is provided in the **SUPPLEMENTARY INFORMATION** section below.

**DATES:** Written comments are due on or before September 9, 2026. This computer matching program will commence on September 9, 2026, and will conclude after 18 months.

**ADDRESSES:** Send comments to Shana Yates, FCC, 45 L Street NE, Washington, DC 20554, or to [Privacy@fcc.gov](mailto:Privacy@fcc.gov).

#### **FOR FURTHER INFORMATION CONTACT:**

Shana Yates at (202) 418-0683 or [Privacy@fcc.gov](mailto:Privacy@fcc.gov).

**SUPPLEMENTARY INFORMATION:** The Lifeline program provides support for discounted broadband and voice services to low-income consumers. Lifeline is administered by the Universal Service Administrative Company (USAC) under FCC direction. Consumers qualify for Lifeline through proof of income or participation in a qualifying program, such as Medicaid, the Supplemental Nutritional Assistance Program (SNAP), Federal Public Housing Assistance, Supplemental Security Income (SSI), Veterans and Survivors Pension Benefit, or various Tribal-specific federal assistance programs.

In a Report and Order adopted on March 31, 2016, (81 FR 33026, May 24, 2016) (*2016 Lifeline Modernization Order*), the Commission ordered USAC to create a National Lifeline Eligibility Verifier ("National Verifier"), including the National Lifeline Eligibility Database (LED), that would match data about Lifeline applicants and subscribers with other data sources to verify the eligibility of an applicant or subscriber. The Commission found that the National Verifier would reduce compliance costs for Lifeline service providers, improve service for Lifeline subscribers, and reduce waste, fraud, and abuse in the program.

#### **Participating Agencies**

Iowa Department of Health and Human Services (source agency); Federal Communications Commission (recipient agency) and Universal Service Administrative Company.

#### **Authority for Conducting the Matching Program**

The authority to conduct the matching program for the FCC's Lifeline program is 47 U.S.C. 254(a) through (c), and (j).

#### **Purpose(s)**

The purpose of this new matching agreement is to verify the eligibility of applicants and subscribers to Lifeline and other Federal programs that use qualification for Lifeline as an eligibility criterion. This new agreement will permit eligibility verification for the Lifeline program by checking an applicant's/subscriber's participation in SNAP in Iowa Department of Health and Human Services. Under FCC rules, consumers receiving these benefits qualify for Lifeline discounts.

### Categories of Individuals

The categories of individuals whose information is involved in the matching program include, but are not limited to, those individuals who have applied for Lifeline; are currently receiving Lifeline; are individuals who enable another individual in their household to qualify for Lifeline; are minors whose status qualifies a parent or guardian for Lifeline; or are individuals who have received Lifeline.

### Categories of Records

The categories of records involved in the matching program include the last four digits of the applicant's Social Security Number, date of birth, first and last name. The National Verifier will transfer these data elements to the Iowa Department of Health and Human Services which will respond either "yes" or "no" that the individual is enrolled in a qualifying assistance program: SNAP administered by the Iowa Department of Health and Human Services.

### System(s) of Records

The records shared as part of this matching program reside in the Lifeline system of records, FCC/WCB-1, Lifeline, which was published in the **Federal Register** at 91 FR 9251 (Feb. 25, 2026).

Federal Communications Commission.

**Marlene Dortch,**  
*Secretary.*

[FR Doc. 2026-16266 Filed 8-7-26; 8:45 am]

**BILLING CODE 6712-01-P**

### GULF COAST ECOSYSTEM RESTORATION COUNCIL

[Docket No.: 108062026-1111-01]

### Notice of Council Updated NEPA Procedures

**AGENCY:** Gulf Coast Ecosystem Restoration Council.

**ACTION:** Notice.

**SUMMARY:** The Gulf Coast Ecosystem Restoration Council (Council) has updated its procedures (NEPA Procedures) for implementing the National Environmental Policy Act (NEPA). The Council's updated NEPA Procedures are available at [www.restorethegulf.gov](http://www.restorethegulf.gov).

**FOR FURTHER INFORMATION CONTACT:** Please send questions by email to John Ettinger by email [john.ettinger@restorethegulf.gov](mailto:john.ettinger@restorethegulf.gov) or telephone (504) 444-3522.

**SUPPLEMENTARY INFORMATION:** On May 5, 2015, the Council issued its NEPA

Procedures (80 FR 25680). The Council's 2015 NEPA Procedures were based on the Council on Environmental Quality (CEQ) NEPA regulations at 40 CFR parts 1500-1508. On January 8, 2026, CEQ issued a final rule removing these regulations (91 FR 618) in accord with Executive Order (E.O.) 14154, *Unleashing American Energy* (January 20, 2025). E.O. 14154 further directed agencies to revise their NEPA implementing procedures consistent with the E.O. Accordingly, the Council has revised its NEPA Procedures. As part of the revision process the Council eliminated a Categorical Exclusion for administrative and routine office activities, which do not meet the definition of major Federal action. The Council's updated NEPA Procedures can be found at [www.restorethegulf.gov](http://www.restorethegulf.gov).

**Keala J. Hughes,**

*Director of External Affairs & Tribal Relations,  
Gulf Coast Ecosystem Restoration Council.*

[FR Doc. 2026-16267 Filed 8-7-26; 8:45 am]

**BILLING CODE 6560-58-P**

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. FDA-2026-P-6298]

### Determination That OZOBAX (Baclofen) Oral Solution, 5 Milligrams/5 Milliliters Was Not Withdrawn From Sale for Reasons of Safety or Effectiveness

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA, Agency, or we) has determined that OZOBAX (baclofen) oral solution, 5 milligrams (mg)/5 milliliters (mL), was not withdrawn from sale for reasons of safety or effectiveness. This determination means that FDA will not begin procedures to withdraw approval of abbreviated new drug applications (ANDAs) that refer to this drug product, and it will allow FDA to continue to approve ANDAs that refer to the product as long as they meet relevant legal and regulatory requirements.

**FOR FURTHER INFORMATION CONTACT:** Heather Dorsey, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 51, Rm. 6269, Silver Spring, MD 20993-0002, 301-796-3600, [heather.dorsey@fda.hhs.gov](mailto:heather.dorsey@fda.hhs.gov).

**SUPPLEMENTARY INFORMATION:** Section 505(j) of the Federal Food, Drug, and

Cosmetic Act (FD&C Act) (21 U.S.C. 355(j)) allows the submission of an ANDA to market a generic version of a previously approved drug product. To obtain approval, the ANDA applicant must show, among other things, that the generic drug product: (1) has the same active ingredient(s), dosage form, route of administration, strength, conditions of use, and (with certain exceptions) labeling as the listed drug, which is a version of the drug that was previously approved, and (2) is bioequivalent to the listed drug. ANDA applicants do not have to repeat the extensive clinical testing otherwise necessary to gain approval of a new drug application (NDA).

Section 505(j)(7) of the FD&C Act requires FDA to publish a list of all approved drugs. FDA publishes this list as part of the "Approved Drug Products With Therapeutic Equivalence Evaluations," which is known generally as the "Orange Book." Under FDA regulations, drugs are removed from the list if the Agency withdraws or suspends approval of the drug's NDA or ANDA for reasons of safety or effectiveness or if FDA determines that the listed drug was withdrawn from sale for reasons of safety or effectiveness (21 CFR 314.162).

A person may petition the Agency to determine, or the Agency may determine on its own initiative, whether a listed drug was withdrawn from sale for reasons of safety or effectiveness. This determination may be made at any time after the drug has been withdrawn from sale, but must be made prior to approving an ANDA that refers to the listed drug (§ 314.161 (21 CFR 314.161)). FDA may not approve an ANDA that does not refer to a listed drug.

OZOBAX (baclofen) oral solution, 5 mg/5 mL, is the subject of NDA 208193, held by Metacel Pharmaceuticals LLC, and initially approved on September 18, 2019. OZOBAX is indicated for the treatment of spasticity resulting from multiple sclerosis, particularly for the relief of flexor spasms and concomitant pain, clonus, and muscular rigidity.

In a letter dated October 13, 2023, Metacel Pharmaceuticals LLC notified FDA that OZOBAX (baclofen) oral solution, 5 mg/5 mL, was being discontinued, and FDA moved the drug product to the "Discontinued Drug Product List" section of the Orange Book.

Novitium Pharma LLC submitted a citizen petition dated June 2, 2026 (Docket No. FDA-2026-P-6298), under 21 CFR 10.30, requesting that the Agency determine whether OZOBAX (baclofen) oral solution, 5 mg/5 mL, was