

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. The Louisiana TIG 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). Please be aware that your entire comment, including your personal identifying information, will become part of the public record. Please note that mailed comments must be postmarked on or before the comment deadline of 30 days following publication of this notice to be considered.

FOR FURTHER INFORMATION CONTACT:

- Louisiana—Joann Hicks, 225–342–5477
- EPA—Douglas Jacobson, 214–665–6692

SUPPLEMENTARY INFORMATION:

Introduction

On April 20, 2010, the mobile offshore drilling unit *Deepwater Horizon*, which was being used to drill a well for BP Exploration and Production, Inc. (BP), in the Macondo prospect (Mississippi Canyon 252–MC252), experienced a significant explosion, fire, and subsequent sinking in the Gulf of Mexico, resulting in the release of an unprecedented volume of oil and other discharges from the rig and from the wellhead on the seabed. The *Deepwater Horizon* oil spill is the largest offshore oil spill in U.S. history, discharging millions of barrels of oil over a period of 87 days. The Trustees conducted the natural resource damage assessment for the *Deepwater Horizon* oil spill under the Oil Pollution Act of 1990 (33 U.S.C. 2701 *et seq.*). Under the OPA, Federal and State agencies act as trustees on behalf of the public to assess natural resource injuries and losses and to determine the actions required to compensate the public for those injuries and losses. The OPA further instructs the designated trustees to develop and implement a plan for the restoration, rehabilitation, replacement, or acquisition of the equivalent of the injured natural resources under their trusteeship, including the loss of use and services from those resources from the time of injury until the time of restoration to baseline (the resource quality and conditions that would exist if the spill had not occurred) is complete.

The *Deepwater Horizon* oil spill Trustees are:

- U.S. Environmental Protection Agency (EPA);
- U.S. Department of the Interior (DOI), as represented by the National Park Service, U.S. Fish and Wildlife Service, and Bureau of Land Management;
- National Oceanic and Atmospheric Administration (NOAA), on behalf of the U.S. Department of Commerce;
- U.S. Department of Agriculture (USDA);
- State of Louisiana Coastal Protection and Restoration Authority (CPRA), Oil Spill Coordinator's Office (LOSCO), Department of Environmental Quality (LDEQ), Department of Wildlife and Fisheries (LDWF), and Department of Natural Resources (LDNR);
- State of Mississippi Department of Environmental Quality;
- State of Alabama Department of Conservation and Natural Resources and Geological Survey of Alabama;
- State of Florida Department of Environmental Protection and Fish and Wildlife Conservation Commission; and
- State of Texas Parks and Wildlife Department, General Land Office, and Commission on Environmental Quality.

On April 4, 2016, the Trustees reached and finalized a settlement of their natural resource damage claims with BP in a Consent Decree approved by the United States District Court for the Eastern District of Louisiana. Pursuant to that Consent Decree, restoration projects in the Louisiana Restoration Area are chosen and managed by the Louisiana TIG. The Louisiana TIG is composed of the following Trustees: CPRA, LOSCO, LDEQ, LDWF, LDNR, EPA, DOI, NOAA, USDA.

Background

The original scope of the Cypremort Improvements project was evaluated in the RP/EA #4, which was published in the **Federal Register** at 83 FR 34571 on July 20, 2018. As proposed in the RP/EA #4, the project would entail a variety of park enhancements including beach restoration, marsh boardwalk and trail construction, road and jetty repairs, and replacement of the breakwater system that helps protect the park's recreational beach. Following completion of the RP/EA #4, the Louisiana Office of State Parks was successful in securing other non-NRDA funding to construct the breakwater system that was originally proposed as a component of the Cypremort Improvements project. The Louisiana TIG prepared the Supplemental RP/EA to evaluate modifications to the Cypremort

Improvements project and consider alternatives, consistent with the purpose and need of the original project. Alternatives considered in the Supplemental RP/EA include replacing the original proposed breakwater system project feature with construction of an RV campground, associated infrastructure, and amenities at the park. The Louisiana TIG prepared the Supplemental RP/EA to inform the public about potential modifications to the Cypremort Improvements project and to seek public comment.

Next Steps

The public is encouraged to review and comment on the Supplemental RP/EA. A public webinar is scheduled to help facilitate the public review and comment process. After the public comment period ends, the Louisiana TIG will consider the comments received before issuing a Final Supplemental RP/EA. A summary of comments received and the Louisiana TIG's responses and any revisions to the document, as appropriate, will be included in the final document. Public comments on the Supplemental RP/EA will inform the Louisiana TIG's decision on whether to select the Cypremort Improvements project, as modified, in the Final Supplemental RP/EA.

Administrative Record

The documents comprising the Administrative Record for the Supplemental RP/EA can be viewed electronically at <https://www.doi.gov/deepwaterhorizon/adminrecord>.

Authority

The authority for this action is the Oil Pollution Act of 1990 (33 U.S.C. 2701 *et seq.*), its implementing NRDA regulations found at 15 CFR part 990, and the NEPA (42 U.S.C. 4321 *et seq.*).

Dated: March 31, 2020.

Benita Best-Wong,

Deputy Assistant Administrator, Office of Water.

[FR Doc. 2020–07263 Filed 4–17–20; 8:45 am]

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FEDERAL COMMUNICATIONS COMMISSION

[3060–1252; FRS 16655]

Information Collection Being Submitted for Review and Approval to Office of Management and Budget

AGENCY: Federal Communications Commission.

ACTION: Notice and request for comments.

SUMMARY: As part of its continuing effort to reduce paperwork burdens, as required by the Paperwork Reduction Act (PRA) of 1995, the Federal Communications Commission (FCC or the Commission) invites the general public and other Federal Agencies to take this opportunity to comment on the following information collection. Pursuant to the Small Business Paperwork Relief Act of 2002, the FCC seeks specific comment on how it might “further reduce the information collection burden for small business concerns with fewer than 25 employees.”

The Commission may not conduct or sponsor a collection of information unless it displays a currently valid Office of Management and Budget (OMB) control number. No person shall be subject to any penalty for failing to comply with a collection of information subject to the PRA that does not display a valid OMB control number.

DATES: Written comments and recommendations for the proposed information collection should be submitted on or before May 20, 2020.

ADDRESSES: Comments should be sent to www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting “Currently under 30-day Review—Open for Public Comments” or by using the search function. Your comment must be submitted into www.reginfo.gov per the above instructions for it to be considered. In addition to submitting in www.reginfo.gov also send a copy of your comment on the proposed information collection to Nicole Ongele, FCC, via email to PRA@fcc.gov and to Nicole.Ongele@fcc.gov. Include in the comments the OMB control number as shown in the **SUPPLEMENTARY INFORMATION** below.

FOR FURTHER INFORMATION CONTACT: For additional information or copies of the information collection, contact Nicole Ongele at (202) 418-2991. To view a copy of this information collection request (ICR) submitted to OMB: (1) Go to the web page <http://www.reginfo.gov/public/do/PRAMain>, (2) look for the section of the web page called “Currently Under Review,” (3) click on the downward-pointing arrow in the “Select Agency” box below the “Currently Under Review” heading, (4) select “Federal Communications Commission” from the list of agencies presented in the “Select Agency” box, (5) click the “Submit” button to the right of the “Select Agency” box, (6) when the list of FCC ICRs currently under review appears, look for the Title of this ICR and then click on the ICR

Reference Number. A copy of the FCC submission to OMB will be displayed.

SUPPLEMENTARY INFORMATION: As part of its continuing effort to reduce paperwork burdens, as required by the Paperwork Reduction Act (PRA) of 1995 (44 U.S.C. 3501–3520), the FCC invited the general public and other Federal Agencies to take this opportunity to comment on the following information collection. Comments are requested concerning: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the Commission, including whether the information shall have practical utility; (b) the accuracy of the Commission’s burden estimates; (c) ways to enhance the quality, utility, and clarity of the information collected; and (d) ways to minimize the burden of the collection of information on the respondents, including the use of automated collection techniques or other forms of information technology. Pursuant to the Small Business Paperwork Relief Act of 2002, Public Law 107–198, see 44 U.S.C. 3506(c)(4), the FCC seeks specific comment on how it might “further reduce the information collection burden for small business concerns with fewer than 25 employees.”

OMB Control Number: 3060–1252.

Title: Application to Participate in Rural Digital Opportunity Fund Auction, FCC Form 183.

Form Number: FCC Form 183.

Type of Review: Revision of a currently approved collection.

Respondents: Business or other for-profit entities, Not-for-profit institutions, and State, Local or Tribal governments.

Number of Respondents and Responses: 500 respondents and 500 responses.

Estimated Time per Response: 7 hours.

Frequency of Response: On occasion reporting requirement.

Obligation to Respond: Required to obtain or retain benefits. Statutory authority for this information collection is contained in 47 U.S.C. 154, 254 and 303(r) of the Communications Act of 1934, as amended.

Estimated Total Annual Burden: 3,500 hours.

Total Annual Costs: No cost.

Nature and Extent of Confidentiality: Although most information collected in FCC Form 183 will be made available for public inspection, the Commission will withhold certain information collected in FCC Form 183 from routine public inspection. Specifically, the Commission will treat certain technical

and financial information submitted in FCC Form 183 as confidential and as though the applicant has requested that this information be treated as confidential trade secrets and/or commercial information. In addition, an applicant may use the abbreviated process under 47 CFR 0.459(a)(4) to request confidential treatment of certain financial information contained in its FCC Form 183 application. However, if a request for public inspection for this technical or financial information is made under 47 CFR 0.461, and the applicant has any objections to disclosure, the applicant will be notified and will be required to justify continued confidential treatment of its request. To the extent that a respondent seeks to have other information collected in FCC Form 183 withheld from public inspection, the respondent may request confidential treatment pursuant to 47 CFR 0.459.

Privacy Act Impact Assessment: No impact(s).

Needs and Uses: The Commission will use the information collected to determine whether applicants are eligible to participate in the Rural Digital Opportunity Fund auction. On January 30, 2020 the Commission adopted the *Rural Digital Opportunity Fund Order*, WC Docket Nos. 19–126, 10–90, FCC 20–5 which will commit up to \$20.4 billion over the next decade to support up to gigabit speed broadband networks in rural America. The funding will be allocated through a multi-round, reverse, descending clock auction that favors faster services with lower latency and encourages intermodal competition in order to ensure that the greatest possible number of Americans will be connected to the best possible networks, all at a competitive cost.

To implement the Rural Digital Opportunity Fund auction, the Commission adopted new rules for the Rural Digital Opportunity Fund auction, including the adoption of a two-stage application process. For the Connect America Fund Phase II auction, applicants that wanted to qualify to bid in the auction were required to submit the FCC Form 183 short-form application. Because the Connect America Fund Phase II auction has ended, the Commission intends to repurpose the FCC Form 183 for the Rural Digital Opportunity Fund auction. Any entity that wishes to participate in the Rural Digital Opportunity Fund auction will be required to submit the FCC Form 183 short-form application to demonstrate its qualifications to bid. Accordingly, the Commission proposes to revise this collection to indicate that it now intends to collect this

information pursuant to section 54.804(a) of the Commission's rules, replacing section 54.315(a) of the Commission's rules. 47 CFR 54.315(a), 54.804(a). The Commission also intends to make several revisions to FCC Form 183, including text changes to reflect the Rural Digital Opportunity Fund auction. Based on the Commission's experience with auctions and consistent with the record, this two-stage collection of information balances the need to collect information essential to conduct a successful auction with administrative efficiency.

Under this information collection, the Commission will collect information that will be used to determine whether an applicant is legally qualified to participate in an auction for Rural Digital Opportunity Fund support. To aid in collecting this information, the Commission will use FCC Form 183, which the public will use to provide the necessary information and certifications. Commission staff will review the information collected on FCC Form 183 as part of the pre-auction process, prior to the start of the auction, and determine whether each applicant satisfies the Commission's requirements to participate in an auction for Rural Digital Opportunity Fund support. Without the information collected on FCC Form 183, the Commission will not be able to determine if an applicant is legally qualified to participate in the auction and has complied with the various applicable regulatory and statutory auction requirements for such participation. This approach is an appropriate assessment of providers for ensuring serious participation without being unduly burdensome.

Federal Communications Commission.

Cecilia Sigmund,

Federal Register Liaison Officer.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Supplemental Evidence and Data Request on Safety of Vaccines Used for Routine Immunization in the United States

AGENCY: Agency for Healthcare Research and Quality (AHRQ), HHS.

ACTION: Request for Supplemental Evidence and Data Submissions

SUMMARY: The Agency for Healthcare Research and Quality (AHRQ) is seeking

scientific information submissions from the public. Scientific information is being solicited to inform our review on *Safety of Vaccines Used for Routine Immunization in the United States*, which is currently being conducted by the AHRQ's Evidence-based Practice Centers (EPC) Program. Access to published and unpublished pertinent scientific information will improve the quality of this review.

DATES: *Submission Deadline* on or before 30 days after the date of this publication in the **Federal Register**.

ADDRESSES:

Email Submissions: epc@ahrq.hhs.gov.

Print Submissions:

Mailing Address: Center for Evidence and Practice Improvement; Agency for Healthcare Research and Quality, ATTN: EPC SEADs Coordinator, 5600 Fishers Lane, Mail Stop 06E53A, Rockville, MD 20857.

Shipping Address (FedEx, UPS, etc.): Center for Evidence and Practice Improvement; Agency for Healthcare Research and Quality, ATTN: EPC SEADs Coordinator, 5600 Fishers Lane, Mail Stop 06E77D, Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT:

Jenae Bennis, Telephone: 301-427-1496 or Email: epc@ahrq.hhs.gov.

SUPPLEMENTARY INFORMATION: The Agency for Healthcare Research and Quality has commissioned the Evidence-based Practice Centers (EPC) Program to complete a review of the evidence for *Safety of Vaccines Used for Routine Immunization in the United States*. AHRQ is conducting this systematic review pursuant to Section 902(a) of the Public Health Service Act, 42 U.S.C. 299a(a).

The EPC Program is dedicated to identifying as many studies as possible that are relevant to the questions for each of its reviews. In order to do so, we are supplementing the usual manual and electronic database searches of the literature by requesting information from the public (e.g., details of studies conducted). We are looking for studies that report on *Safety of Vaccines Used for Routine Immunization in the United States*, including those that describe adverse events. The entire research protocol is available online at: <https://effectivehealthcare.ahrq.gov/products/safety-vaccines/protocol>.

This is to notify the public that the EPC Program would find the following information on *Safety of Vaccines Used for Routine Immunization in the United States* helpful:

- A list of completed studies that your organization has sponsored for this

indication. In the list, please *indicate whether results are available on ClinicalTrials.gov along with the ClinicalTrials.gov trial number*.

- *For completed studies that do not have results on ClinicalTrials.gov*, a summary, including the following elements: study number, study period, design, methodology, indication and diagnosis, proper use instructions, inclusion and exclusion criteria, primary and secondary outcomes, baseline characteristics, number of patients screened/eligible/enrolled/lost to follow-up/withdrawn/analyzed, effectiveness/efficacy, and safety results.

- *A list of ongoing studies that your organization has sponsored for this indication*. In the list, please provide the *ClinicalTrials.gov* trial number or, if the trial is not registered, the protocol for the study including a study number, the study period, design, methodology, indication and diagnosis, proper use instructions, inclusion and exclusion criteria, and primary and secondary outcomes.

- Description of whether the above studies constitute *ALL Phase II and above clinical trials* sponsored by your organization for this indication and an index outlining the relevant information in each submitted file.

Your contribution is very beneficial to the Program. Materials submitted must be publicly available or able to be made public. Materials that are considered confidential; marketing materials; study types not included in the review; or information on indications not included in the review cannot be used by the EPC Program. This is a voluntary request for information, and all costs for complying with this request must be borne by the submitter.

The draft of this review will be posted on AHRQ's EPC Program website and available for public comment for a period of 4 weeks. If you would like to be notified when the draft is posted, please sign up for the email list at: <https://www.effectivehealthcare.ahrq.gov/email-updates>.

The systematic review will answer the following questions. This information is provided as background. AHRQ is not requesting that the public provide answers to these questions.

Key Questions (KQ)

KQ 1: What is the evidence that vaccines included in the immunization schedule recommended for adults in the United States (<https://www.cdc.gov/vaccines/schedules/hcp/imz/adult.html>) are safe in the short term (within 42 days following immunization) or long term (>42 days after immunization)?