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: http://www.epa.gov/compliance/nepa/eisdata.html.

**EIS No. 20130381, Draft EIS, FHWA, TX, US 181 Harbor Bridge Project, Comment Period Ends: 03/03/2014, Contact: Gregory Punskie 512–536–5960.**

**EIS No. 20130382, Draft EIS, NPS, 00, Glen Canyon National Recreation Area Off-road Vehicle Management Plan, Comment Period Ends: 03/04/2014, Contact: Brian Carey 928–608–6209.**

**EIS No. 20130383, Final EIS, USACE, CA, Berryessa Creek Element Coyote and Berryessa Creek California Flood Control Project, Review Period Ends: 02/03/2014, Contact: Tyler Stalker 916–557–5107.**

**Amended Notices**

**EIS No. 20130325, Draft EIS, NPS, MO, Ozark National Scenic Riverways Draft General Management Plan, Wilderness Study, Comment Period Ends: 02/07/2014, Contact: William Black 573–323–4236, Revision to the FR Notice Published 12/06/2013; Extending Comment Period from 01/08/2014 to 02/07/2014.**

**EIS No. 20130360, Final EIS, USFS, AZ, Rosemont Copper Project, Proposed Construction, Operation with Concurrent Reclamation and Closure of an Open-Pit Copper Mine, Review Period Ends: 02/14/2014, Contact: Mindy Vogel 520–388–8300, Revision to FR Notice Published 12/20/2013; Correcting the Review Period End Date to read 02/14/2014. Dated: December 30, 2013.**

**Dawn Roberts, Management Analyst, NEPA Compliance Division, Office of Federal Activities.**

**ENVIRONMENTAL PROTECTION AGENCY**


**Petition To Add the Oil and Gas Extraction Industry, Standard Industrial Classification Code 13, to the List of Facilities Required To Report Under the Toxics Release Inventory; Notice of Receipt of Petition**

**AGENCY:** Environmental Protection Agency (EPA).  

**ACTION:** Notice of receipt of petition.

**SUMMARY:** The Environmental Integrity Project (EIP) and sixteen other organizations submitted a petition to the Environmental Protection Agency (EPA), dated October 24, 2012, requesting that EPA add the Oil and Gas Extraction sector, Standard Industrial Classification (SIC) code 13, to the scope of sectors covered by the Toxics Release Inventory (TRI) under section 313 of the Emergency Planning and Community Right-To-Know Act (EPCRA). The petition also requests that EPA publish the petition in the Federal Register. This Federal Register Notice provides notice of receipt of this petition, along with the Docket Identification Number that can be used to view the petition and related documents. EPA is not soliciting public comment regarding this notice.

**FOR FURTHER INFORMATION CONTACT:** Gilbert Mears, Toxics Release Inventory Program Division, Office of Environmental Information (mail code 2844T), Environmental Protection Agency, 1200 Pennsylvania Avenue NW., Washington, DC 20460; telephone number: (202) 566–0954; fax number: (202) 566–0715; email address: mears.gilbert@epa.gov.

**SUPPLEMENTARY INFORMATION:**

**I. General Information**

**A. How can I get copies of this document and other related information?**

1. Docket. EPA has established a docket for this action under Docket ID No EPA–HQ–TRI–2013–0281. Publicly available docket materials are available either electronically through www.regulations.gov or in hard copy at the “Petition to Add the Oil and Gas Extraction Industry, Standard Industrial Classification Code 13, to the List of Facilities Required To Report under the Toxics Release Inventory” Docket in the EPA Docket Center, (EPA/DC) EPA West, Room 3334, 1301 Constitution Ave. NW., Washington, DC. The EPA Docket Center Public Reading Room is open from 8:30 a.m. to 4:30 p.m., Monday through Friday, excluding legal holidays. The telephone number for the Public Reading Room is (202) 566–1744, and the telephone number for the “Petition to Add the Oil and Gas Extraction Industry, Standard Industrial Classification Code 13, to the List of Facilities Required To Report under the Toxics Release Inventory” Docket is (202) 566–1752.


Arnold E. Layne,  
Director, Office of Information Analysis and Access, Office of Environmental Information.

**[FR Doc. 2013–31484 Filed 1–2–14; 8:45 am]**

**BILLING CODE 6560–50–P**

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Medicare Program; Appellant Forum Regarding the Administrative Law Judge Hearing Program for Medicare Claim Appeals**

**AGENCY:** Office of Medicare Hearings and Appeals (OMHA), HHS.

**ACTION:** Notice of Meeting.

**SUMMARY:** This notice announces an Office of Medicare Hearings and Appeals (OMHA) Medicare Appellant Forum. The purpose of this event is to provide updates to OMHA appellants on the status of OMHA operations; to relay information on a number of OMHA initiatives designed to mitigate a growing backlog in the processing of Medicare appeals at the OMHA-level of the administrative appeals process; and provide information on measures that appellants can take to make the administrative appeals process work more efficiently at the OMHA-level.

**DATES:** Meeting Date: The OMHA Medicare Appellant Forum announced in this notice will be held on Wednesday, February 12, 2014. The OMHA Medicare Appellant Forum will begin at 10:00 a.m. Eastern Standard Time (e.s.t.) and check-in will begin at 9:00 a.m. e.s.t.

**Deadline for Registration of Attendees and Requests for Special Accommodation:** The deadline to register to attend the OMHA Medicare Appellants Forum and request a special accommodation, as provided for in the American’s with Disabilities Act, is 5:00
The Office of Medicare Hearings and Appeals (OMHA), a staff division within the Office of the Secretary of the U.S. Department of Health and Human Services (HHS), administers the nationwide Administrative Law Judge hearing program for Medicare claim and entitlement appeals, and has established the Office of Medicare Hearings and Appeals (OMHA). OMHA was established in July 2005 pursuant to section 931 of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (Pub. L. 108–173), which required the transfer of responsibility for the Administrative Law Judge hearing level of the Medicare claim and entitlement appeals process from the Social Security Administration to HHS. OMHA was established to improve service to appellants and reduce the average 368-day time frame to receive a hearing decision that appellants experienced with the Social Security Administration, in accordance with 90-day adjudication time frame mandated by the Medicare, Medicaid, and SCHIP Benefits Improvement and Protection Act of 2000 (Pub. L. 106–554).

At the time of OMHA’s establishment, it was envisioned that OMHA would receive the traditional claim and entitlement appeals workload from the Medicare Part A and Part B programs, and the Medicare Part C Medicare Advantage program; and a new workload of appeals from the Medicare Part D Prescription Drug program and the Income Related Monthly Adjustment Amount (IRMMA) premium surcharges assessed by the Social Security Administration. However, beginning in fiscal year 2010, new workloads including permanent establishment of the Recovery Audit (RA) program and termination of several demonstration projects involving Medicare State Agencies (MSA), have emerged that had not been built into the OMHA workload models. The steady growth in traditional Medicare appeals combined with these new workloads has strained OMHA’s ability to meet the adjudication time frame mandate while maintaining quality.

As a result of the anticipated workload increase from the traditional appeals and the increased workload resulting from MSA appeals and the RA program, a backlog of appeals began to form in fiscal year 2012 in which more requests for hearing were filed than could be adjudicated. In 2013, appeals pending in the RA program grew to over 136,000, further exacerbating the backlog of cases and resulting in a substantial increase in the adjudication time frame.

As the unprecedented growth in claim appeals continues to exceed the available adjudication resources to address appeals, OMHA is taking measures to mitigate the workload increase and planning future activities to bring efficiencies to the appeals process at the OMHA level. One of the immediate measures has been to ensure that the relatively small numbers of beneficiary-initiated appeals are being immediately addressed by prioritizing their cases. For the remaining cases, OMHA has suspended assignments of new requests for hearing until an adjudicator becomes available, which will allow cases to be assigned more efficiently on a first in/first out basis as an Administrative Law Judge’s case docket is able to accommodate additional workload. In addition, OMHA is vigorously pursuing an electronic case adjudication processing environment (ECAPE) to bring new efficiencies and appellant-access to the OMHA-level of the appeals process.

The OMHA Medicare Appellant Forum will address these initiatives, as well as potential future initiatives, and will solicit input from the appellant community to help OMHA evaluate its policies and procedures to achieve meaningful backlog reduction strategies and process efficiencies while remaining compliant with applicable legal authorities.

II. Medicare Claim Appeal Appellant Forum and Conference Calling/Webinar Information

A. Format of the OMHA Medicare Appellant Forum

As noted in section I. of this notice, OMHA is conducting this outreach to appellants of the Medicare claim appeals process to provide updates on initiatives to mitigate a growing backlog in the processing Medicare appeals at the OMHA level, and to solicit input to achieve meaningful backlog reduction strategies and process efficiencies. Information regarding the OMHA Medicare Appellant Forum can be found on the OMHA Web site at: http://www.hhs.gov/omha/index.html.

The majority of the forum will be reserved for presentations of workload data, processes and policy discussions, and recommendations from agency presenters. The time for each presentation will be approximately 30 to 60 minutes and will be based on the material being addressed in the presentation.

Questions and comments from in-person attendees will be solicited at the
end of each planned presentation and during a separate question and answer session as time permits. In addition, questions related to the OMHA-level of the Medicare claim appeals process will also be accepted on the attendee registration, for potential response during the appropriate presentation.

B. Conference Call, Live Streaming, and Webinar Information

For participants who cannot attend the OMHA Medicare Appellant Forum in person, there may be an option to view and participate in the OMHA Medicare Appellant Forum via live streaming technology and/or a webinar. Information on the whether these capabilities will be available as part of this forum will be posted on the OMHA Web site at: http://www.hhs.gov/omha/index.html. Please continue to check the Web site for updates on this upcoming event.

Disclaimer: We cannot guarantee reliability for live streaming technology and/or a webinar.

III. Registration Instructions

The OMHA Executive Office is coordinating attendee registration for the OMHA Medicare Appellant Forum. While there is no registration fee, individuals planning to attend the forum must register to attend. In-person participation is limited to two (2) representatives from each organization. Additional individuals can participate by telephone conference or webinar if these services are made available. Information on participation by telephone conference or webinar will be posted on the OMHA Web site at: http://www.hhs.gov/omha/index.html. Registration may be completed online at the following web address: http://www.hhs.gov/omha/index.html. Seating capacity for in-person attendees is limited to the first 400 registrants.

After completing the registration, online registrants will receive a confirmation email which they should bring with them to the meeting(s). If you are unable to register online, you may register by calling the OMHA Appellant Forum line at 202-356-3000 or by sending an email to OSOMHAAppellantForum@hhs.gov. Please include your first and last name, title, organization, address, office telephone number, and email address. If seating capacity has been reached, you will be notified that the meeting has reached capacity.

IV. Security, Building, and Parking Guidelines

Because the OMHA Medicare Appellant Forum will be conducted on Federal property, for security reasons, any persons wishing to attend these meetings must register by the date specified in the DATES section of this notice. Please allow sufficient time to go through the security checkpoints. It is suggested that you arrive at the Wilbur J. Cohen building, located at 330 Independence Ave. SW., Washington, DC 20024, no later than 9:30 a.m. e.s.t. if you are attending the forum in person so that you will be able to arrive promptly for the meeting.

Security measures include the following:
- Presentation of photographic identification to the Federal Protective Service or Guard Service personnel.
- Passing through a metal detector and inspection of items brought into the building. We note that all items brought to the Cohen Building, whether personal or for the purpose of demonstration or to support a demonstration, are subject to inspection. We cannot assume responsibility for coordinating the receipt, transfer, transport, storage, set-up, safety, or timely arrival of any personal belongings or items used for demonstration or to support a demonstration.

Note: Individuals who are not registered in advance will not be permitted to enter the building and will be unable to attend the forum in person. The public may not enter the building earlier than 45 minutes prior to the convening of the forum.

Attendees must enter the Cohen Building thru the C-Street entrance and proceed to the registration desk. All visitors must be escorted in areas other than the auditorium area and access to the rest rooms on the same level in the building. Seating capacity is limited to the first 400 registrants.

Parking in Federal buildings is not available for this event. In addition, street side and commercial parking is extremely limited in the downtown area. Attendees are advised to use Metro-rail to either the Federal Center SW station (Blue/Orange line) or the L’Enfant Plaza station (Yellow/Green or Blue/Orange lines). The Wilbur J. Cohen building is approximately 1 ½ blocks from each of these Metro-rail stops.

(DEPARTMENT OF HEALTH AND HUMAN SERVICES)

Centers for Disease Control and Prevention

[30Day–14–0892]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and Budget (OMB) in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call (404) 639–7570 or send an email to omb@cdc.gov. Send written comments to CDC Desk Officer, Office of Management and Budget, Washington, DC 20503 or by fax to [202] 395–5806. Written comments should be received within 30 days of this notice.

Proposed Project
Clostridium difficile Infection (CDI) Surveillance

Steady increases in the rate and severity of Clostridium difficile infection (CDI) indicate a clear need to conduct longitudinal assessments to continue to monitor changes in CDI epidemiology, including changes in risk factors for disease, as well as increases in incidence and severity of illness related to this pathogen.

The title and the goals of the project have remained the same since the publication of the 60-day Federal Register Notice and there were no changes in burden estimates or data collection forms from what is shown in the current inventory.

The surveillance population will consist of persons residing in the catchment area of the participating Emerging Infections Program (EIP) sites who are 1 year of age or older. This surveillance poses no more than minimal risk to the study participants as there will be no interventions or modifications to the care study participants receive.

EIP surveillance personnel will perform active case finding from laboratory reports of stool specimens testing positive for C. difficile toxin and abstract data on cases using a standardized case report form. For a subset of cases (e.g., community-associated C. difficile cases) sites will administer a health interview.