order for the LRN Program Office to determine the ability of the Network to respond to a biological or chemical threat event. The sensitivity of all information associated with the LRN requires the LRN Program Office to obtain personal information about all individuals accessing the LRN Web site. In addition, the LRN Program Office must be able to contact all laboratory personnel during an event so each laboratory staff member that obtains access to the restricted LRN Web site must provide his or her contact information to the LRN Program Office.

As a requirement of membership, LRN Laboratories must report all biological and chemical testing results to the LRN Program at CDC using a CDC developed software tool called the LRN Results Messenger. This information is essential for surveillance of anomalies, to support response to an event that may involve multiple agencies and to manage limited resources. LRN Laboratories must also participate in and report results for Proficiency Testing Challenges or Validation Studies. LRN Laboratories participate in multiple Proficiency Testing Challenges, Exercises and/or Validation Studies every year consisting of five to 500 simulated samples provided by the LRN Program Office. It is necessary to conduct such challenges in order to verify the testing capability of the LRN Laboratories.

The rarity of biological or chemical agents perceived to be of bioterrorism concern prevent some LRN Laboratories from maintaining proficiency as a result of day-to-day testing. Simulated samples are therefore distributed to ensure proficiency across the LRN. The results obtained from testing these simulated samples must also be entered into Results Messenger for evaluation by the LRN Program Office.

During a surge event resulting from a bioterrorism or chemical terrorism attack, LRN Laboratories are also required to submit all testing results using LRN Results Messenger. The LRN Program Office requires these results in order to track the progression of a bioterrorism event and respond in the most efficient and effective way possible and for data sharing with other Federal partners involved in the response. The number of samples tested during a response to a possible event could range from 10,000 to more than 500,000 samples depending on the length and breadth of the event. Since there is potentially a large range in the number of samples for a surge event, CDC estimates the annualized burden for this event will be 2,250,000 hours or 625 responses per respondent.

Semiannually the LRN Program Office may conduct a Special Data Call to obtain additional information from LRN Member Laboratories in regards to biological or chemical terrorism preparedness. Special Data Calls are conducted using the LRN Web site. There is no cost to the respondents other than their time.

### ESTIMATED ANNUALIZED BURDEN HOURS

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hrs)</th>
<th>Total burden (in hrs)</th>
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<tr>
<td>Public Health Laboratories</td>
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<td>625</td>
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<td>2,250,000</td>
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<td>Special Data Call</td>
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<td>10</td>
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<td>Total</td>
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<td>2,385,300</td>
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</table>


Ron Otten,
Director, Office of Science Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2012–31182 Filed 1–2–13; 8:45 am]
BILLING CODE 4162–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30 Day–13–0696]

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


Background and Brief Description

CDC is requesting a 3-year approval for revision to the previously approved project. The purpose of this revision is to continue collecting standardized HIV prevention program evaluation data from health departments and community-based organizations (CBOs) who receive federal funds for HIV prevention activities. Grantees have the option of key-entering or uploading data to a CDC-provided web-based software application (EvaluationWeb®).

The following changes have occurred since project 0920–0696 has been implemented: (1) The previous reporting system (PEMS) has been replaced by a more efficient reporting software. (2) Many data variables that were previously required or optional but reported have been deleted in order to reduce data reporting burden on grantees. Other variables have been added or modified to adapt to changes in HIV prevention and the National HIV/AIDS Strategic Plan. (3) Reporting has been changed from quarterly to semiannual. (4) The number of grantees has changed as new FOAs were awarded.
The evaluation and reporting process is necessary to ensure that CDC receives standardized, accurate, thorough evaluation data from both health departments and CBO grantees. For these reasons, CDC developed standardized NHM&E variables through extensive consultation with representatives from health departments, CBOs, and national partners (e.g., The National Alliance of State and Territorial AIDS Directors, Urban Coalition of HIV/AIDS Prevention Services, and National Minority AIDS Council).

CDC requires CBOs and health departments who receive federal funds for HIV prevention to report non-identifying, client-level and aggregate-level, standardized evaluation data to: (1) Accurately determine the extent to which HIV prevention efforts are carried out, what types of agencies are providing services, what resources are allocated to those services, to whom services are being provided, and how these efforts have contributed to a reduction in HIV transmission; (2) improve ease of reporting to better meet these data needs; and (3) be accountable to stakeholders by informing them of HIV prevention activities and use of funds in HIV prevention nationwide.

**Estimated Annualized Burden Hours**

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
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<td>H/RR Data</td>
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<td>67</td>
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<td></td>
<td>HIV Testing Data</td>
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<td>NHM&amp;E Data Training</td>
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<td>2</td>
<td>30/60</td>
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<tr>
<td></td>
<td>H/RR Data</td>
<td>200</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>NHM&amp;E Data Training</td>
<td>200</td>
<td>2</td>
<td>20</td>
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</table>


Ron A. Otten,
Director, Office of Scientific Integrity (OSI), Office of the Associate Director for Science (OADS), Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2012–31599 Filed 1–2–13; 8:45 am]
BILLING CODE 4163–18–P

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Medicare & Medicaid Services**

**[CMS–3278–NC]**

**Medicare Program: Request for Information on Hospital and Vendor Readiness for Electronic Health Records Hospital Inpatient Quality Data Reporting**

**AGENCY:** Centers for Medicare & Medicaid Services (CMS), HHS.

**ACTION:** Request for information.

**SUMMARY:** This document is a request for information from hospitals, electronic health record (EHR) vendors, and other interested parties regarding hospital readiness beginning calendar year 2014 discharges to electronically report certain patient-level data under the Hospital Inpatient Quality Reporting (IQR) Program using the Quality Reporting Document Architecture (QRDA) Category I.

**DATES:** The information solicited in this document must be received at the address provided below, no later than 5 p.m. eastern standard time (e.s.t.) on January 22, 2013.

**ADDRESSES:** In commenting, refer to file code CMS–3278–NC. Because of staff and resource limitations, we cannot accept comments by facsimile (FAX) transmission.

You may submit comments in one of four ways (please choose only one of the ways listed):

1. **Electronically.** You may submit electronic comments on this regulation to http://www.regulations.gov. Follow the “Submit a comment” instructions.

2. **By regular mail.** You may mail written comments to the following address only: Centers for Medicare & Medicaid Services, Department of Health and Human Services, Attention: CMS–3278–NC, P.O. Box 8013, Baltimore, MD 21244–8013.

   Please allow sufficient time for mailed comments to be received before the close of the comment period.

3. **By express or overnight mail.** You may send written comments to the following address only: Centers for Medicare & Medicaid Services, Department of Health and Human Services, Attention: CMS–3278–NC, Mail Stop C4–26–05, 7500 Security Boulevard, Baltimore, MD 21244–1850.

4. **By hand or courier.** Alternatively, you may deliver (by hand or courier) your written comments ONLY to the following addresses:


   (Because access to the interior of the Hubert H. Humphrey Building is not readily available to persons without Federal government identification, commenters are encouraged to leave their comments in the CMS drop slots located in the main lobby of the building. A stamp-in clock is available for persons wishing to retain a proof of filing by stamping in and retaining an extra copy of the comments being filed.)

   b. For delivery in Baltimore, MD—Centers for Medicare & Medicaid Services, Department of Health and Human Services, 7500 Security Boulevard, Baltimore, MD 21244–1850.

   If you intend to deliver your comments to the Baltimore address, call telephone number (410) 786–9994 in advance to schedule your arrival with one of our staff members.

   Comments erroneously mailed to the addresses indicated as appropriate for hand or courier delivery may be delayed and received after the comment period.