

warrantors would now disclose this information even if there were no statute or rule requiring them to do so, staff's estimates likely overstate the PRA-related burden attributable to the Rule. Moreover, the Warranty Rule has been in effect since 1976, and warrantors have long since modified their warranties to include the information the Rule requires.

Based on conversations with various warrantors' representatives over the years, staff has concluded that eight hours per year is a reasonable estimate of warrantors' PRA-related burden attributable to the Warranty Rule.⁵ This estimate takes into account ensuring that new warranties and changes to existing warranties comply with the Rule. Based on recent Census data, staff now estimates that there are 15,922 manufacturers covered by the Rule.⁶ This results in an annual burden estimate of approximately 127,376 hours (15,922 manufacturers × 8 hours of burden per year).

Total annual labor costs: \$16,941,000, rounded to the nearest thousand.

Labor costs are derived by applying appropriate hourly cost figures to the burden hours described above. The work required to comply with the Warranty Rule—ensuring that new warranties and changes to existing warranties comply with the Rule—requires a mix of legal analysis and clerical support. Staff estimates that half of the total burden hours (63,688 hours) requires legal analysis at an average hourly wage of \$250 for legal professionals,⁷ resulting in a labor cost of \$15,922,000. Assuming that the remaining half of the total burden hours requires clerical work at an average hourly wage of \$16, the resulting labor cost is approximately \$1,019,008. Thus, the total annual labor cost is approximately \$16,941,008 (\$15,922,000 for legal professionals + \$1,019,008 for clerical workers).

Total annual capital or other nonlabor costs: \$0.

The Rule imposes no appreciable current capital or start-up costs. As stated above, warrantors have already

⁵ FTC staff recently contacted two manufacturing associations—the Association of Home Appliance Manufacturers and the National Association of Manufacturers—but we have not received any additional information that further clarifies this estimate.

⁶ Because some manufacturers likely make products that are not priced above \$15 or not intended for household use—and thus would not be subject to the Rule—this figure is likely an overstatement.

⁷ Staff has derived an hourly wage rate for legal professionals based upon industry knowledge. The clerical wage rate used in this Notice is based on recent data from the Bureau of Labor Statistics National Compensation Survey.

modified their warranties to include the information the Rule requires. Rule compliance does not require the use of any capital goods, other than ordinary office equipment, which providers would already have available for general business use.

Willard K. Tom,
General Counsel.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

5th Annual PHEMCE Stakeholders Workshop and BARDA Industry Day

AGENCY: Department of Health and Human Services, Office of the Secretary.

ACTION: Notice.

SUMMARY: The Department of Health and Human Services (HHS) is pleased to announce the upcoming 5th Annual Public Health Emergency Medical Countermeasures Enterprise (PHEMCE) Stakeholders Workshop and BARDA Industry Day to be held January 10–12, 2011 at the Walter E. Washington Convention Center in Washington, DC. This annual PHEMCE event will bring together private- and public-sector stakeholders including: Federal Officials, International Governments, Industry, Healthcare Providers, First Responders, Community-Based Organizations, and other interested audiences. Attendees will have opportunities to participate in Medical Countermeasure focused forums on:

- Pre-Event Positioning of Medical Countermeasures.
- Emergency Planning for Vulnerable Populations.
- Industry Feedback on Contracting Issue.
- Medical Countermeasures Development: Expanding the Pipeline and Exploring Multi-Use Potential.
- BARDA Industry Day Presentations.

This free Workshop will also address current state of public health emergency medical countermeasure preparedness plans and opportunities to enhance national response capabilities. BARDA Industry Day provides a unique opportunity for biotechnology and pharmaceutical industry representatives to showcase their latest advances in vaccines, therapeutics, diagnostics, and platform technologies targeting chemical, biological, radiological, nuclear, and naturally emerging threats, including pandemic influenza.

DATES: The 5th Annual PHEMCE Stakeholders Workshop and BARDA Industry Day will be held January 10–12, 2011. Each day will begin at 9 a.m.

ADDRESSES: The Workshop will be held at the Walter E. Washington Convention Center, 801 Mount Vernon Place, NW., Washington, DC 20001.

Registration: There is no fee to attend; however, space is limited and registration is required. Registration and the preliminary agenda are available online at: <http://www.medicalcountermeasures.gov>.

FOR FURTHER INFORMATION CONTACT: L. Paige Rogers, Office of Policy and Planning, Office of the Assistant Secretary for Preparedness and Response at 330 Independence Ave., SW., Room G640, Washington, DC 20201, e-mail at BARDA@hhs.gov, or by phone at 202-260-0365.

Dated: September 16, 2010.

Nicole Lurie,

Assistant Secretary for Preparedness and Response.

[FR Doc. 2010-26047 Filed 10-14-10; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

[Document Identifier: OS-0990-New; 30-day notice]

Agency Information Collection Request; 30-Day Public Comment Request

AGENCY: Office of the Secretary, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Office of the Secretary (OS), Department of Health and Human Services, is publishing the following summary of a proposed collection for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency's functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, e-mail your request, including your address, phone number,

OMB number, and OS document identifier, to *Sherette.funncoleman@hhs.gov*, or call the Reports Clearance Office on (202) 690-5683. Send written comments and recommendations for the proposed information collections within 30 days of this notice directly to the OS OMB Desk Officer; faxed to OMB at 202-395-5806.

Proposed Project: ONC State HIE Performance Measures and Progress Report—OMB No. 0990-NEW—Office

of the National Coordinator for Health Information Technology.

Abstract: The purpose of the State Health Information Exchange Cooperative Agreement Program, as authorized by Section 3013 of the American Recovery and Reinvestment Act is to provide grants to States and Qualified State Designated Entities is to facilitate and expand the secure, electronic movement and use of health information among organizations according to national recognized

standards. As part of that project, States and Qualified State Designated Entities are required to provide biannual program progress reports and report on performance measures during the implementation phase of the cooperative agreement. This request is for those two data gathering requirements. The data collection will last four years, which is the duration of the project, and this request is for the data collection for the first three years of that project period.

ESTIMATED ANNUALIZED BURDEN TABLE

Forms (if necessary)	Type of respondent	Number of respondents	Number of responses per respondent	Average burden per response (in hrs.)	Total burden hours
Evaluation performance measures ...	State government or Qualified State Designated Entity.	56	2	175	19,600
Program progress report	State government or Qualified State Designated Entity.	56	2	8	896
Total	20,496

Terry Nicolosi,
 Director, Office of Resources Management;
 Office of the Chief Information Officer.
 [FR Doc. 2010-25917 Filed 10-14-10; 8:45 am]
 BILLING CODE 4150-45-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

[Document Identifier: OS-0990-NEW; 30-day notice]

Agency Information Collection Request; 30-Day Public Comment Request

AGENCY: Office of the Secretary, HHS.
 In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Office of the Secretary (OS), Department of Health and Human Services, is publishing the following summary of a proposed collection for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed

information collection for the proper performance of the agency's functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, e-mail your request, including your address, phone number, OMB number, and OS document identifier, to *Sherette.funncoleman@hhs.gov*, or call the Reports Clearance Office on (202) 690-5683. Send written comments and recommendations for the proposed information collections within 30 days of this notice directly to the OS OMB Desk Officer; faxed to OMB at 202-395-5806.

Proposed Project: ONC State HIE State Plans—OMB No. 0990-NEW—Office of the National Coordinator for Health Information Technology.

Abstract: The purpose of the State Health Information Exchange Cooperative Agreement Program, as authorized by Section 3013 of the American Recovery and Reinvestment Act is to provide grants to States and Qualified State Designated Entities is to facilitate and expand the secure, electronic movement and use of health information among organizations according to national recognized standards. Section 3013 requires States and Qualified State Designated Entities to have approved State Plans, consisting of strategic and operational components, before funding can be used for implementation activities. The State Plans must be submitted to the National Coordinator for Health Information Technology during the first year of the project period in order to receive implementation funding through the cooperative agreement. Annual updates to the State plans will be required in the three remaining project periods. The data collection will last four years, which is the duration of the project, and this request is for the data collection for the first three years of that project period.

ESTIMATED ANNUALIZED BURDEN TABLE

Forms (if necessary)	Type of respondent	Number of respondents	Number of responses per respondent	Average burden per response (in hrs.)	Total burden hours
State Plans (Strategic and Operational).	State Government or Qualified State Designated Entity.	56	1	10,024	561,244