TOTAL ESTIMATED ANNUALIZED BURDEN—HOURS

Information collection (see above list "Burden statement" for legend)	Number of Respondents	Number of responses per respondent	Average burden per response (hours)	Total burden hours
(a)	63	1	8	504
(b)	63	1	12	756
(c)	63	3	12	2268
(d)	63	3	14	2646
(e)	63	1	10	630
(f)	63	1	8	504
(g)	63	3	8	1512
(h)	63	3	3	567
(i)	63	3	8	1512
Total	63	19	83	10,899

Darius Taylor,

Deputy, Information Collection Clearance Officer.

[FR Doc. 2014–04931 Filed 3–5–14; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Clinical Laboratory Improvement Advisory Committee, Centers for Disease Control and Prevention: Notice of Charter Renewal

This gives notice under the Federal Advisory Committee Act (Public Law 92–463) of October 6, 1972, that the Clinical Laboratory Improvement Advisory Committee, Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS), has been renewed for a 2-year period through February 19, 2016.

For information, contact Devery Howerton, Ph.D., Designated Federal Officer, Clinical Laboratory Improvement Advisory Committee, 1600 Clifton Road NE., Mailstop E–56, Atlanta, Georgia 30333, telephone 404–498–2602 or via email at dxh7@cdc.gov.

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Elaine L. Baker,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2014–04935 Filed 3–5–14; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2014-N-0237]

Report to Congress; Report on the Food and Drug Administration's Policy To Be Proposed Regarding Premarket Notification Requirements for Modifications to Legally Marketed Devices; Notice to Public of Web Site Location of Report to Congress

AGENCY: Food and Drug Administration, HHS

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or the Agency) is announcing the Web site location where the Agency has posted the report entitled "Report to Congress; Report on FDA's Policy to be Proposed Regarding Premarket Notification Requirements for Modifications to Legally Marketed Devices." In addition, FDA has established a docket where stakeholders may provide comments.

DATES: Submit either electronic or written comments by June 4, 2014. **ADDRESSES:** Submit electronic comments on this document to *http://www.regulations.gov.* Submit written comments to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT:

Mike Ryan, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, Rm. 1615, Silver Spring, MD 20993–0002, 301–796–6283.

SUPPLEMENTARY INFORMATION:

I. Background

The Food and Drug Administration Safety and Innovation Act (FDASIA) (Pub. L. 112–144) became law on July 9,

2012. FDASIA added section 510(n)(2) to the Federal Food, Drug, and Cosmetic Act (the FD&C Act) (21 U.S.C. 360(n)(2)). This new provision requires, no later than 18 months after enactment of FDASIA, the Secretary of Health and Human Services to submit to the Committee on Energy and Commerce of the House of Representatives and the Committee on Health, Education, Labor, and Pensions of the Senate a report on when a premarket notification under section 510(k) of the FD&C Act (or a "510(k)") should be submitted for a modification to a legally marketed 510(k) device. This report fulfills that requirement.

This notice announces the Web site location of "Report to Congress; Report on FDA's Policy to be Proposed Regarding Premarket Notification Requirements for Modifications to Legally Marketed Devices." FDA invites interested persons to submit comments on this report. FDA has established a docket where comments may be submitted (see ADDRESSES). FDA believes this docket is an important tool for receiving information from interested parties and for sharing this information with the public. To access "Report to Congress; Report on FDA's Policy to be Proposed Regarding Premarket Notification Requirements for Modifications to Legally Marketed Devices," visit FDA's Web site http:// www.fda.gov/AboutFDA/CentersOffices/ OfficeofMedicalProductsandTobacco/ CDRH/CDRHReports/ucm269873.htm.

II. Request for Comments

Interested persons may submit either electronic comments regarding this document to http://www.regulations.gov or written comments to the Division of Dockets Management (see ADDRESSES). It is only necessary to send one set of comments. Identify comments with the docket number found in brackets in the heading of this document. Received