above-noted motor vehicle dealers. This resulted in an estimate of 248 motor vehicle dealers subject to the FTC’s jurisdiction. After deducting the latter figure from the total of 750 respondents, 502 respondents were left to divide 50:50 between the agencies. With rounding, the FTC apportioned 251 of those respondents to its burden estimates; adding to that the estimated total of 248 motor vehicle dealers resulted in 499 respondents for the FTC.

B. FTC Share of Burden Hours: 998 Hours

Staff assumed that respondents will each spend approximately 2 hours to monitor compliance with the Rule. Thus, 499 respondents for the FTC multiplied by the two hour estimate per respondent resulted in 998 burden hours apportioned to the FTC.

C. FTC Share of Labor Costs: $249,500

Staff assumed that in-house legal counsel for respondents would handle most of the compliance review, and at an estimated average hourly wage of $250 per hour.

D. Capital/Non-Labor Costs: $0

Assumption: Capital and other nonlabor costs should be minimal, at most, since the Rule has been in effect several years, with covered entities now equipped to provide the required notice.

Based on staff’s review of industry data and its experience in this area, we have no information to suggest that these figures are not still valid.

Request for Comments

You can file a comment online or on paper. Write “Prescreen Opt-Out Disclosure Rule: FTC File No. P075417” on your comment. Your comment—including your name and your state—will be placed on the public record of this proceeding, including, to the extent practicable, on the public Commission Web site, at http://www.ftc.gov/os/publiccomments.shtm. As a matter of discretion, the Commission tries to remove individuals’ home contact information from comments before placing them on the Commission Web site.

Because your comment will be made public, you are solely responsible for making sure that your comment does not include any sensitive personal information, such as a Social Security number, date of birth, driver’s license number or other state identification number or foreign country equivalent, passport number, financial account number, or credit or debit card number. You are also solely responsible for making sure that your comment does not include any sensitive health information, such as medical records or other individually identifiable health information. In addition, do not include any “[t]rade secret or any commercial or financial information which is privileged or confidential,” as discussed in Section 6(f) of the FTC Act, 15 U.S.C. 46(f), and FTC Rule 4.10(a)(2). 16 CFR 4.10(a)(2). In particular, do not include competitively sensitive information such as costs, sales statistics, inventories, formulas, patterns, devices, manufacturing processes, or customer names.

If you want the Commission to give your comment confidential treatment, you must file it in paper form, with a request for confidential treatment, and you must follow the procedure explained in FTC Rule 4.9(c), 16 CFR 4.9(c). Your comment will be kept confidential only if the FTC General Counsel, in his or her sole discretion, grants your request in accordance with the law and the public interest. Postal mail addressed to the Commission is subject to delay due to heightened security screening. As a result, the Commission encourages you to submit your comments online. To make sure that the Commission considers your online comment, you must file it at https://ftcpublic.commentworks.com/ftc/prescreenoptoutpro by following the instructions on the web-based form. If this Notice appears at http://www.regulations.gov, you also may file a comment through that Web site.

If you file your comment on paper, write “Prescreen Opt-Out Disclosure Rule: FTC File No. P075417” on your comment and on the envelope, and mail it to the following address: Federal Trade Commission, Office of the Secretary, 600 Pennsylvania Avenue NW., Suite CC–5610, (Annex J), Washington, DC 20580, or deliver your comment to the following address: Federal Trade Commission, Office of the Secretary, Constitution Center, 400 7th Street SW., 5th Floor, Suite 5610, (Annex J), Washington, DC 20024. If possible, submit your paper comment to the Commission by courier or overnight service.

The FTC Act and other laws that the Commission administers permit the collection of public comments to consider and use in this proceeding as appropriate. The Commission will consider all timely and responsive public comments that it receives on or before July 25, 2016. You can find more information, including routine uses permitted by the Privacy Act, in the Commission’s privacy policy, at http://www.ftc.gov/ftc/privacy.htm.

David C. Shonka, Acting General Counsel.

[FR Doc. 2016–12330 Filed 5–24–16; 8:45 am]
BILLING CODE 6750–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS–10418]

Agency Information Collection Activities: Submission for OMB Review; Comment Request

ACTION: Notice.

SUMMARY: The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS’ intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (PRA), federal agencies are required to publish a notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, and to allow a second opportunity for public comment on the notice. Interested persons are invited to send comments regarding the 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.

DATES: Comments on the collection(s) of information must be received by the OMB desk officer by June 24, 2016.

ADDRESSES: When commenting on the proposed information collections, please reference the document identifier or OMB control number. To be assured consideration, comments and
recommendations must be received by the OMB desk officer via one of the following transmissions:

OMB, Office of Information and Regulatory Affairs, Attention: CMS Desk Officer, Fax Number: (202) 395–5806 or Email: OIRA_omb.eop.gov

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of following:
2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to PaperworkReductionActof1995.
3. Call the Reports Clearance Office at (410) 786–1326.

FOR FURTHER INFORMATION CONTACT:
Reports Clearance Office at (410) 786–1326.

SUPPLEMENTARY INFORMATION:
Under the Paperwork Reduction Act of 1995 (PRA) (44 U.S.C. 3501–3520), federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. The term “collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires federal agencies to publish a 30-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice that summarizes the following proposed collection(s) of information for public comment:

1. Type of Information Collection Request: Revision of currently approved collection; Title of Information Collection: Annual Medical Loss Ratio (MLR) and Rebate Calculation Report and MLR Rebate Notices: Use: Under section 2718 of the Affordable Care Act and implementing regulation at 45 CFR part 158, a health insurance issuer (issuer) offering group or individual health insurance coverage must submit a report to the Secretary concerning the amount the issuer spends each year on claims, quality improvement expenses, non-claims costs, Federal and State taxes and licensing and regulatory fees, the amount of earned premium, and beginning with the 2014 reporting year, the amounts related to the reinsurance, risk corridors, and risk adjustment programs established under sections 1341, 1342, and 1343, respectively, of the Affordable Care Act. An issuer must provide an annual rebate if the amount it spends on certain costs compared to its premium revenue (excluding Federal and State taxes and licensing and regulatory fees) does not meet a certain ratio, referred to as the medical loss ratio (MLR). Each issuer is required to submit annually MLR data, including information about any rebates it must provide, on a form prescribed by CMS, for each State in which the issuer conducts business. Each issuer is also required to provide a rebate notice to each policyholder that is owed a rebate and each subscriber of policyholders that are owed a rebate for any given MLR reporting year. Additionally, each issuer is required to maintain for a period of seven years all documents, records and other evidence that support the data included in each issuer’s annual report to the Secretary. Under section 1342 of the Patient Protection and Affordable Care Act and implementing regulation at 45 CFR part 153, issuers of qualified health plans (QHPs) must participate in a risk corridors program. A QHP issuer will pay risk corridors charges or be eligible to receive payments based on the ratio of the issuer’s allowable costs to the target amount. Each QHP issuer is required to submit an annual report to CMS concerning the issuer’s allowable costs, allowable administrative costs, premium, and proportion of market premium in QHPs. Risk corridors premium information that is specific to an issuer’s QHPs is collected through a separate plan-level data form, which is included in this information collection. CMS received a total of 3 public comments on a number of specific issues regarding the notice of the revised MLR PRA package. CMS has taken into consideration all of the comments and has modified the information collection instruments and instructions (the 2015 MLR Annual Reporting Form and Instructions and the 2015 Risk Corridors Plan-Level Data Form and Instructions) in order to correct minor errors and to provide additional clarifications. Form Number: CMS–10418 (OMB control number: 0938–1164); Frequency: Annually; Affected Public: Private Sector (Business or other for-profits and not-for-profit institutions); Number of Respondents: 538; Number of Burden Hours: 181; Total Annual Hours: 235,148. (For policy questions regarding this collection contact Christina Whitefield at (301) 492–4172.)

Dated: May 18, 2016.

William N. Parham, III,
Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2016–12085 Filed 5–24–16; 8:45 am] BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Health Resources and Services Administration

Ryan White HIV/AIDS Program Part C
HIV Early Intervention Services Program Existing Geographic Service Area

AGENCY: Health Resources and Services Administration, HHS.
ACTION: Notice of Class Deviation from Competition Requirements for Ryan White HIV/AIDS Program (RWHAP) Part C HIV Early Intervention Services Program Existing Geographic Service Area (EISEGA).

SUMMARY: The HIV/AIDS Bureau (HAB) is requesting a class deviation from the competition requirements in order to provide one-year extensions with funds to 346 Ryan White HIV/AIDS Program (RWHAP) Part C HIV Early Intervention Services Program Existing Geographic Service Area (EISEGA) recipients. The purpose of the Part C EISEGA program is to provide HIV primary care in the outpatient setting to targeted low income, underinsured people living with HIV. HAB is finalizing an evaluation of the Part C EISEGA program and development of a new data-driven methodology. This methodology is aimed at ensuring that awards are based on a consistent approach to promote a rational and sustainable allocation of resources while ensuring responsiveness to geographic and healthcare financing considerations, indicators of need, and results along the HIV care continuum. HAB expects to re-compete the entire program in fiscal year (FY) 2018. One-year extensions with funds for all 346 Part C EISEGA recipients enables HAB to finalize the evaluation and methodology development and engage recipients and relevant stakeholders with regard to this new approach prior to implementation and without disrupting the provision of critical HIV primary medical care services to the current RWHAP clients served by these recipients. Pending the availability of funds, the amount of each FY 2017 award will be based on a