

difficult to do so within the period of time allowed by this notice, you should advise the contact listed below as soon as possible.

ADDRESSES: Direct all PRA comments to Cathy Williams, FCC, via email *PRA@fcc.gov* and to *Cathy.Williams@fcc.gov*.

FOR FURTHER INFORMATION CONTACT: For additional information about the information collection, contact Cathy Williams at (202) 418-2918.

SUPPLEMENTARY INFORMATION:

OMB Control No.: 3060-1029.

Title: Data Network Identification Code (DNIC).

Form No.: N/A.

Type of Review: Extension of a currently approved collection.

Respondents: Business or other for-profit.

Number of Respondents: 5 respondents; 5 responses.

Estimated Time per Response: .25 hours.

Frequency of Response: On occasion reporting requirement.

Obligation to Respond: Required to obtain or retain benefits. The statutory authority for this collection is contained in 47 U.S.C. 151, 154(i)-(j), 201-205, 211, 214, 219, 220, 303(r), 309 and 403.

Total Annual Burden: 1 hour.

Annual Cost Burden: N/A.

Privacy Act Impact Assessment: N/A.

Nature and Extent of Confidentiality:

In general, there is no need for confidentiality with this collection of information.

Needs and Uses: This collection will be submitted as an extension (no change in reporting or recordkeeping requirements) after this 60-day comment period to Office of Management and Budget (OMB) in order to obtain the full three year clearance.

A Data Network Identification Code (DNIC) is a unique, four-digit number designed to provide discrete identification of individual public data networks. The DNIC is intended to identify and permit automated switching of data traffic to particular networks. The FCC grants the DNICs to operators of public datanetworks on an international protocol. The operators of public data networks file an application for a DNIC on the Internet-based, International Bureau Filing System (IBFS). The DNIC is obtained free of charge on a one-time only basis unless there is a change in ownership or the owner chooses to relinquish the code to the FCC. The Commission's lack of an assignment of DNICs to operators of public data networks would result in technical problems that prevent the identification and automated switching of data traffic to particular networks.

Federal Communications Commission.

Sheryl D. Todd,

Deputy Secretary, Office of the Secretary.

[FR Doc. 2015-23544 Filed 9-18-15; 8:45 am]

BILLING CODE 6712-01-P

FEDERAL ELECTION COMMISSION

Sunshine Act Meetings

AGENCY: Federal Election Commission.

DATE AND TIME: Thursday, September 17, 2015 at 10:00 a.m.

PLACE: 999 E Street NW., Washington, DC (Ninth Floor).

STATUS: This Meeting Will Be Open To The Public.

Federal Register Notice of Previous Announcement—80 FR 55115

This item was also discussed:

Second Motion to Set Priorities and Scheduling on Pending Enforcement Matters Awaiting Reason-to-Believe Consideration

Individuals who plan to attend and require special assistance, such as sign language interpretation or other reasonable accommodations, should contact Shawn Woodhead Werth, Secretary and Clerk, at (202)694-1040, at least 72 hours prior to the meeting date.

PERSON TO CONTACT FOR INFORMATION:

Judith Ingram, Press Officer, Telephone: (202) 694-1220.

Shelley E. Garr,

Deputy Secretary of the Commission.

[FR Doc. 2015-24047 Filed 9-17-15; 4:15 pm]

BILLING CODE 6715-01-P

FEDERAL RESERVE SYSTEM

Change in Bank Control Notices; Acquisitions of Shares of a Bank or Bank Holding Company

The notificants listed below have applied under the Change in Bank Control Act (12 U.S.C. 1817(j)) and § 225.41 of the Board's Regulation Y (12 CFR 225.41) to acquire shares of a bank or bank holding company. The factors that are considered in acting on the notices are set forth in paragraph 7 of the Act (12 U.S.C. 1817(j)(7)).

The notices are available for immediate inspection at the Federal Reserve Bank indicated. The notices also will be available for inspection at the offices of the Board of Governors. Interested persons may express their views in writing to the Reserve Bank indicated for that notice or to the offices of the Board of Governors. Comments

must be received not later than October 6, 2015.

A. Federal Reserve Bank of Minneapolis (Jacquelyn K. Brunmeier, Assistant Vice President) 90 Hennepin Avenue, Minneapolis, Minnesota 55480-0291:

1. *Marian Dahlgren, and Lee Dahlgren, individually, Greg Dahlgren, and Lee Dahlgren, as co-trustees of the Marian Dahlgren Trust*, all of Vergas, Minnesota; to join the Dahlgren Family group and acquire voting shares of Vergas Bancorporation, Inc., and thereby indirectly acquire voting shares of Vergas State Bank, both in Vergas, Minnesota.

Board of Governors of the Federal Reserve System, September 16, 2015.

Michael J. Lewandowski,

Associate Secretary of the Board.

[FR Doc. 2015-23584 Filed 9-18-15; 8:45 am]

BILLING CODE 6210-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Agency for Healthcare Research and Quality, HHS.

ACTION: Notice.

SUMMARY: This notice announces the intention of the Agency for Healthcare Research and Quality (AHRQ) to request that the Office of Management and Budget (OMB) approve the proposed information collection project: "*Online Submission Form for Supplemental Evidence and Data for Systematic reviews for the Evidence-based Practice Center Program.*" In accordance with the Paperwork Reduction Act, 44 U.S.C. 3501-3521, AHRQ invites the public to comment on this proposed information collection.

DATES: Comments on this notice must be received by November 20, 2015.

ADDRESSES: Written comments should be submitted to: Doris Lefkowitz, Reports Clearance Officer, AHRQ, by email at *doris.lefkowitz@AHRQ.hhs.gov*.

Copies of the proposed collection plans, data collection instruments, and specific details on the estimated burden can be obtained from the AHRQ Reports Clearance Officer.

FOR FURTHER INFORMATION CONTACT: Doris Lefkowitz, AHRQ Reports Clearance Officer, (301) 427-1477, or by email at *doris.lefkowitz@AHRQ.hhs.gov*.

SUPPLEMENTARY INFORMATION:**Proposed Project**

This is a new activity of AHRQ's Evidence-based Practice Center Program.

Evidence-based Practice Center Program

AHRQ's Evidence-based Practice Center (EPC) Program develops evidence reports and technology assessments on topics relevant to clinical and other health care organization and delivery issues—specifically those that are common, expensive, and/or significant for the Medicare and Medicaid populations. For example recent reviews have focused on clinical conditions, such as “Treatment of Nonmetastatic Muscle-Invasive Bladder Cancer”; health delivery topics such as “Management Strategies to Reduce Psychiatric Admissions”; and specific technologies such as “Imaging Techniques for Treatment Evaluation for Metastatic Breast Cancer.” These evidence reports include systematic reviews and technical briefs, and provide an essential foundation from which to understand what we know from existing research and what critical research gaps remain. These reports, reviews, and technology assessments are based on rigorous, comprehensive syntheses and analyses of the scientific literature on topics. EPC reports and assessments emphasize explicit and detailed documentation of methods, rationale, and assumptions. EPC reports are conducted in accordance with an established policy on financial and nonfinancial interests. These scientific syntheses may include meta-analyses and cost analyses.

The EPC Program supports AHRQ's mission by synthesizing and disseminating the available research as a “science partner” with private and public organizations in their efforts to improve the quality, effectiveness, and appropriateness of health care. The EPC Program is a trusted source of rigorous, comprehensive, and unbiased evidence reviews for stakeholders. The resulting evidence reports and technology assessments are used by Federal and State agencies, private-sector professional societies, health delivery systems, providers, payers, and others committed to evidence-based health care. These end users may use EPC Program evidence reports to inform policy decisions, clinical practice guidelines, and other health care decisions.

EPC research has the following goals:

- Use research methods to gather knowledge on the effectiveness of

certain treatments for specific medical conditions, both published and unpublished, to evaluate the quality of research studies and the evidence from these studies.

- Promote the use of evidence in health care decision making to improve health care and health.

- Identify research gaps to inform future research investments.

The Institute of Medicine standards for quality systematic reviews include an assessment of publication bias through the identification of unpublished studies. This is an important source for bias which could affect the nature and direction of research findings. Identifying and including the results of these additional unpublished studies may provide a more complete and accurate assessment of an intervention's effect on outcomes. An important way to identify unpublished studies is through requests to medical device manufacturers, pharmaceutical companies, and other intervention developers.

The proposed project involves sending a request letter to relevant medical device manufacturers, pharmaceutical companies and other intervention developers to invite them to submit unpublished studies or other scientific information to the EPC Program Web site, with one request per systematic review topic. Because research on each topic must be completed in a timely manner in order for it to be useful, the collections are never ongoing—there is one request and collection per topic. Investigators in the EPC Program will review the information and assess potential risk of bias from both published and unpublished studies and its impact on the EPC Program's findings. AHRQ believes the display of these assessments in the systematic review's evidence tables will improve the response and submission rates of industry stakeholders by informing the health care community of the impact of potential bias on the research conclusions, and for health care decision making.

This activity is being conducted by AHRQ's EPC Program through its contractor, the Scientific Resource Center (SRC), pursuant to AHRQ's statutory authority to conduct and support research on health care and on systems for the delivery of such care and to disseminate government-funded research relevant to comparative clinical effectiveness research. 42 U.S.C. 299a(a); 42 U.S.C. 299b–37(a).

Method of Collection

To achieve the goals of this project the following data collections will be implemented:

- Online Submission Form Instrument. This information is collected for the purposes of providing supplemental evidence and data for systematic reviews (SEADS). The online submission form (OSF) collects data from respondents on their organization name, their product's name, and whether they are providing all information on requested studies characteristic of the review in progress. This happens following receipt of a request letter from the SRC. These requests will be sent to relevant sponsors of preventative and treatment interventions (e.g., medical device manufacturers, pharmaceuticals, and other intervention and health care system developers), with one request per topic. For the purposes of meta-analyses, trial summary data from missing and unidentified studies are sought. For the purposes of constructing evidence tables and quality ratings (e.g. on public reporting of cost measures or health information exchange), data can vary (e.g., URLs, study designs, and consumer-mediated exchange forms). Information on both completed and ongoing studies are requested.

The EPC Program, through the SRC, currently uses a **Federal Register** notice and broad-based email announcement to stakeholders to allow the public to know about each topic, and the opportunity to submit scientific information. In 2014, the Program sent 517 notifications to 336 industry stakeholders. Of those 517 announcements sent, 14.1% received a response; 56.2% of the responses (or 7.9% of all requests) contained submissions of information on the results of interventions. This experience has prompted this proposed project.

The additional use of direct requests to relevant organizations would improve the Program's ability to obtain this information. Contacting intervention sponsors for missing and potentially unidentified studies could improve the impact of research efforts and downstream dissemination efforts and could positively impact the health of individuals, burdened by poor health along with their supporting communities. Including information about response data to these requests to more accurately characterize the completeness of the evidence in the systematic reviews may also address this issue.

The proposed project does not duplicate other available sources of this

information. Available study registries and databases may not be complete to sufficiently inform the Program’s research.

The purpose of SEADS requests is not to collect generalizable data, but to supplement the published and grey literature searches EPC investigators are conducting. Furthermore, considering the evidence and data included in responses collected from industry stakeholders, an assessment pertaining to the completeness of the evidence-base will be produced. This, AHRQ believes, will increase the value of AHRQ’s research reviews to end users and potentially provide stakeholders a

better understanding of how their submissions are used.

Estimated Annual Respondent Burden

Exhibit 1 presents estimates of the reporting burden hours for the data collection efforts. Time estimates are based on pilot testing of materials and what can reasonably be requested of respondents. The number of respondents listed in “Number of respondents per SEADS request” of Exhibit 1 reflects a projected 80% response rate.

Online Submission Form: A form for submitting scientific evidence and data related to medical interventions

sponsored by organizations and individuals such as pharmaceutical companies and independent researchers. The form has three required fields: The organization’s name, the intervention in question, and whether the information they provide is all the information they know to exist. They may upload documents and they are also provided a data entry form if they wish to offer greater details on their studies.

An Optional Data Entry Form is available as an alternative to the Online Submission form. The time requirements for response would be same as the Online Submission Form.

EXHIBIT 1—ESTIMATED ANNUALIZED BURDEN HOURS

Form name	Number of respondents per SEADS request	Number of responses per respondent	Hours per response	Total burden hours per SEADS
Online Submission Form (OSF)	70	1	15/60	17.5
Total	70	1	15/60	17.5

EXHIBIT 2—ESTIMATED ANNUALIZED COST HOURS

Form name	Number of SEADS requests	Total burden hours per SEADS	Average hourly wage rate*	Total cost burden
OSF	70	17.5	\$55.48 ^a	\$970.90
Total	70	17.5	55.48	970.90

*Occupational Employment Statistics, May 2014 National Occupational Employment and Wage Estimates United States, U.S. Department of Labor, Bureau of Labor Statistics. http://www.bls.gov/oes/current/oes_nat.htm#b29-0000.

^aBased on the mean wages for *Public Relations and Fundraising Managers, 11–2031*, the occupational group most likely tasked with completing the OSF.

Request for Comments

In accordance with the Paperwork Reduction Act, comments on AHRQ’s information collection are requested with regard to any of the following: (a) Whether the proposed collection of information is necessary for the proper performance of AHRQ health care research and health care information dissemination functions, including whether the information will have practical utility; (b) the accuracy of AHRQ’s estimate of burden (including hours and costs) of the proposed collection(s) of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information upon the respondents, including the use of automated collection techniques or other forms of information technology.

Comments submitted in response to this notice will be summarized and included in the Agency’s subsequent request for OMB approval of the proposed information collection. All

comments will become a matter of public record.

Sharon Arnold,

Deputy Director,

[FR Doc. 2015–23573 Filed 9–18–15; 8:45 am]

BILLING CODE 4160–90–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Notice of Meetings

AGENCY: Agency for Healthcare Research and Quality (AHRQ), HHS.

ACTION: Notice of Five AHRQ Subcommittee Meetings.

SUMMARY: The subcommittees listed below are part of AHRQ’s Health Services Research Initial Review Group Committee. Grant applications are to be reviewed and discussed at these meetings. Each subcommittee meeting will commence in open session before

closing to the public for the duration of the meeting. These meetings will be closed to the public in accordance with 5 U.S.C. App. 2 section 10(d), 5 U.S.C. 552b(c)(4), and 5 U.S.C. 552b(c)(6).

DATES: See below for dates of meetings:

1. *Healthcare Effectiveness and Outcomes Research (HEOR)*
Date: October 7, 2015 (Open from 8:30 a.m. to 9:00 a.m. on October 7th and closed for remainder of the meeting)
2. *Health System and Value Research (HSVR)*
Date: October 7, 2015 (Open from 8:30 a.m. to 9:00 a.m. on October 7th and closed for remainder of the meeting)
3. *Healthcare Safety and Quality Improvement Research (HSQR)*
Date: October 14–15, 2015 (Open from 8:00 a.m. to 8:30 a.m. on October 14th and closed for remainder of the meeting)
4. *Health Care Research and Training (HCRT)*
Date: October 15–16, 2015 (Open from 8:00 a.m. to 8:30 a.m. on October 15th and closed for remainder of the meeting)