

identified by the Director of the Federal Emergency Management Administration as being subject to special flood hazards. The Riegle Community Development Act requires that each institution must also provide a copy of the notice to the servicer of the loan (if different from the originating lender).

Dated: March 18, 2002.

Federal Deposit Insurance Corporation.

Robert E. Feldman,

Executive Secretary.

[FR Doc. 02-6951 Filed 3-21-02; 8:45 am]

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FEDERAL ELECTION COMMISSION

[Notice 2002-4]

The Voting System Standards and an Opportunity to Publicly Voice Previously Submitted Comments

AGENCY: Federal Election Commission.

ACTION: Notice of public hearing.

SUMMARY: The Federal Election Commission is announcing a public hearing on the December 13, 2001, release of the Voting System Standards.

DATES: The hearing will be held at 10:00 a.m. on Wednesday, April 17, 2002. All requests to testify must be received by the Commission by April 7, 2002. Requests to testify are limited to election officials, members of the National Association of State Election Directors' Voting System Standards Board, and those parties who have previously submitted written comments to the June 16, 2001, and/or December 13, 2001, release of the Voting System Standards.

ADDRESSES: Requests to testify should be addressed to Penelope Bonsall, Director of the Office of Election Administration, and must be submitted in either written or electronic form. Due to recent delays in mail service to government offices, electronic or fax submissions are encouraged to ensure timeliness. Written requests to testify should be sent to the Federal Election Commission, 999 E Street, NW., Washington, DC 20463. Faxed requests should be sent to (202) 219-8500, with printed copy follow-up to insure legibility. Electronic mail requests should be sent to vss@fec.gov. Persons sending requests by electronic mail must include their full name, electronic mail address and postal service address within the text of the request.

Commission hearings are held in the Commission's ninth floor meeting room, 999 E Street, NW., Washington, DC.

FOR FURTHER INFORMATION CONTACT: Penelope Bonsall, Director of the Office

of Election Administration, 999 E Street, NW., Washington, DC 20463, (202) 694-1095 or (800) 424-9530, ext. 1095.

SUPPLEMENTARY INFORMATION: The Voting System Standards (the "Standards") were originally promulgated in 1990. Technological and commercial innovations during the last decade have demanded that the Standards be updated, and the project to revise them was begun in 1998. The revised Standards have two volumes. Volume I provides functional and technical requirements for a number of system types and configurations. Volume II provides testing specifications for the requirements in Volume I. Both Volumes are available at the Commission's web site (<http://www.fec.gov/pages/vss/vss.html>). The Commission previously released for public comment a draft of the first volume on June 16, 2001. 66 FR 35978. During this comment period, the Commission received 38 sets of comments from 39 parties.

Subsequently, the Commission released the entire draft Standards on December 13, 2001. 66 FR 65708. The comment period for the December 13, 2001, draft release ended on February 1, 2002. FR Notice 2001. Twenty-seven sets of comments from twenty-three parties were received by the Commission in response to the December 13, 2001, release. Four commenters requested to testify at a public hearing if one is held.

After considering these requests and the other comments received to date in response to the notice, the Commission believes a public hearing would be helpful in considering the issues raised by the draft Standards. The hearing will be held at 10:00 a.m. on April 17, 2002.

Dated: March 18, 2002.

David M. Mason,

Chairman, Federal Election Commission.

[FR Doc. 02-6948 Filed 3-21-02; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-02-30]

Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the Centers for Disease Control and

Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call the CDC Reports Clearance Officer on (404) 498-1210.

Comments are invited on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection 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 on respondents, including through the use of automated collection techniques or other forms of information technology. Send comments to Anne O'Connor, CDC Assistant Reports Clearance Officer, 1600 Clifton Road, MS-D24, Atlanta, GA 30333. Written comments should be received within 60 days of this notice.

Proposed Project: Outcome Evaluation of HIV Prevention Programs Focusing on Prevention Case Management Interventions Implemented by the Directly-funded Community-Based Organizations (CBOs)—New—National Center for HIV, STD and Tuberculosis Prevention (NCHSTP), Centers for Disease Control and Prevention (CDC). This evaluation is necessary to understand the impact of CDC's expenditures and efforts to support CBOs and for modifying and improving the HIV prevention case management efforts of CBOs. This data collection will provide standardized data and allow CDC to (a) assess the implementation and effectiveness of HIV prevention case management (PCM) interventions through process and outcome evaluations; (b) determine the degree of adherence to the CBOs' documented HIV PCM intervention protocol, and through quality assurance efforts, to revise program implementation as necessary; (c) understand the behavioral impact of these programs; and (d) provide useful information for CBO program planners and implementers.

Three CBOs funded under Program Announcement 01000, Community-Based Strategies to Increase HIV Testing of Persons at High Risk in Communities of Color, successfully competed for additional funds from Program Announcement 01159, Outcome Evaluation of HIV Prevention Programs with a focus on Prevention Case Management Interventions and Group-Level Interventions Implemented by CDC's Directly-funded Community-

Based Organizations, to conduct an outcome evaluation of their PCM interventions for two years. These CBOs administer baseline social-behavioral questionnaires as part of program services. Each CBO will report on the

PCM program that it has implemented, and, as part of the research project, will conduct two short follow-up social-behavioral questionnaires with clients to assess changes in participant risk behaviors. Incentives will be given to

CBO respondents to complete follow-up assessments. This is a two-year project; each of the three CBOs is estimated to collect data from 100 clients each year. There are no costs to respondents.

Respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
CBO Clients (year—1)	300	1	30/60	150
CBO Clients (year—2)	300	1	30/60	150
Total				300

Dated: March 18, 2002.

Nancy Cheal,

Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-02-31]

Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call the CDC Reports Clearance Officer on (404) 498-1210.

Comments are invited on: (a) Whether the proposed collection of information

is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection 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 on respondents, including through the use of automated collection techniques or other forms of information technology. Send comments to Anne O'Connor, CDC Assistant Reports Clearance Officer, 1600 Clifton Road, MS-D24, Atlanta, GA 30333. Written comments should be received within 60 days of this notice.

Proposed Project: Breast, Colorectal, and Prostate Cancer Patterns of Care, Reoccurrence, and Survival (CBOs)—New—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC). Invasive cancers of the breast, colon and rectum, and prostate impose a substantial burden of disease in the United States (U.S.) and are expected to account for approximately 42 percent of the estimated 1.3 million invasive cancers that will be diagnosed during 2002. Breast and colorectal cancers are particularly of high public health

importance because of current widespread activities in place for early diagnosis and treatment.

Even though these cancers are of high public importance, statewide central cancer registries are not likely to capture complete follow-up information or detailed information on treatment modalities other than surgery. Also, data on extent of disease at diagnosis are often limited. In order to expand the uses of their data to include survival and patterns of care studies and clinical research, registries may need to collect additional information. Through re-abstracting representative samples of cases from population-based, central cancer registries from 1997, this pattern of care study will assess the quality of stage and treatment data. Estimates of the proportions of patients who received the standard of care for localized breast, localized prostate, and stage III colon cancers will be determined as well. Registries participating in the study will send data to the CDC for some analyses. Data for the patterns of care study and for the CONCORD Study, a collaborative project between the CDC and cancer registries in the U.S. and Europe, will be re-abstracted from medical records at the same time. The annualized estimated cost to respondents is \$2,056,000.

Respondents	Number of respondents	Number of responses/re-spondent	Average burden/response (in hours)	Total burden (in hours)
Physicians (M.D., D.O.)	4440	1	15/60	1,110
Total				1,110