

Board, are available for immediate inspection at the Federal Reserve Bank indicated. The applications also will be available for inspection at the offices of the Board of Governors. Interested persons may express their views in writing on the standards enumerated in the BHC Act (12 U.S.C. 1842(c)). If the proposal also involves the acquisition of a nonbanking company, the review also includes whether the acquisition of the nonbanking company complies with the standards in section 4 of the BHC Act (12 U.S.C. 1843). Unless otherwise noted, nonbanking activities will be conducted throughout the United States. Additional information on all bank holding companies may be obtained from the National Information Center website at [www.ffiec.gov/nic/](http://www.ffiec.gov/nic/).

Unless otherwise noted, comments regarding each of these applications must be received at the Reserve Bank indicated or the offices of the Board of Governors not later than March 17, 2008.

**A. Federal Reserve Bank of Kansas City** (Todd Offenbacher, Assistant Vice President) 925 Grand Avenue, Kansas City, Missouri 64198-0001:

1. *Prime Bank Group, Inc.*, to become a bank holding company by acquiring 100 percent of the voting shares of Prime Bank (in organization), both in Edmond, Oklahoma.

**B. Federal Reserve Bank of Dallas** (W. Arthur Tribble, Vice President) 2200 North Pearl Street, Dallas, Texas 75201-2272:

1. *National Bank & Trust Employee Stock Ownership Plan With 401(k) Provisions*; to become a bank holding company by acquiring up to 26 percent of the voting shares of First La Grange Bancshares, Inc., and indirectly acquire voting shares of National Bank & Trust, all of La Grange, Texas.

Board of Governors of the Federal Reserve System, February 15, 2008.

**Robert deV. Frierson,**

*Deputy Secretary of the Board.*

[FR Doc. E8-3202 Filed 2-20-08; 8:45 am]

BILLING CODE 6210-01-S

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Agency for Healthcare Research and Quality

#### Office for Civil Rights: Audio Conference on Proposed Regulations Related to Patient Safety

**AGENCY:** Agency for Healthcare Research and Quality, HHS; Office for Civil Rights, HHS.

**ACTION:** Notice of audio conference.

**SUMMARY:** The U.S. Department of Health and Human Services' Agency for Healthcare Research and Quality Director Dr. Carolyn Clancy and Office for Civil Rights Deputy Director of Health Information Privacy Susan McAndrew will host a joint audio conference February 29, 2008 from 2-3 p.m. (Eastern Standard Time) to discuss the recently published proposed regulation regarding Patient Safety and Quality Improvement and statutory confidentiality protections. The purpose of this audio conference is to facilitate public understanding of the proposed regulation and rulemaking process outlined in the Notice of Proposed Rulemaking published in the **Federal Register** February 12, 2008. To register for the audio conference, log on to <http://www.academyhealth.org/ahrq/psaudio/>.

**DATES:** The live audio conference will be Feb. 29 from 2-3 p.m. (Eastern Standard Time).

**ADDRESSES:** The proposed regulation can be viewed on the Federal eRulemaking Portal at <http://www.regulations.gov/fdmspublic/ContentViewer?objectId=09000064803acce8&disposition=attachment&contentType=html>.

The audio conference is open to everyone; however, discussions during this forum will not be included in official public comments.

Public comment on the proposed regulations will be accepted through April 14, 2008.

Comments can be submitted by any of the following methods: Federal eRulemaking Portal: <http://www.regulations.gov/fdmspublic/component/main?main=SubmitComment&o=09000064803acce8>.

Comments should include the agency name (Agency for Healthcare Research and Quality and/or Office for Civil Rights) and RIN 0919-AA01.

Mail/Hand Delivery/Courier: Center for Quality Improvement and Patient Safety, Attention: Patient Safety Act Notice of Proposed Rulemaking Comments, Agency for Healthcare Research and Quality, 540 Gaither Road, Rockville, Maryland 20850.

Comments sent by facsimile (FAX) transmission or electronic mail will not be accepted.

Comments received through the eRulemaking Portal can be viewed online at either of the Web sites listed above. All comments received through the eRulemaking Portal, mail, and hand delivery/courier are available for public inspection at the AHRQ Information Resources Center, which is located at

540 Gaither Road, Rockville, Maryland 20850. The Information Resources Center is open from 8:30 a.m. to 5 p.m. Eastern Standard Time, Monday through Friday.

**FOR FURTHER INFORMATION CONTACT:** Ellen Crown, Agency for Healthcare Research and Quality, 301-427-1258 or [ellen.crown@ahrq.hhs.gov](mailto:ellen.crown@ahrq.hhs.gov).

**SUPPLEMENTARY INFORMATION:** Under the Patient Safety and Quality Improvement Act of 2005 (Patient Safety Act), the Secretary is authorized to list Patient Safety Organizations (PSOs), organizations that will work with providers to collect and analyze patient safety related data. The Statute sets forth and the recently published proposed regulation explains certifications that must be submitted by entities in order to be listed as PSOs. PSOs will provide analysis of data and feedback to providers to assist them in improving patient safety.

The Patient Safety Act protects the confidentiality of data shared by providers prepared by the PSO as well as other related materials, defined in the statute and proposed regulations. This legal protection of information addresses significant barriers that currently exists—the fear of legal liability or sanctions that can result from reporting a patient safety event. Strong confidentiality provisions are key to encouraging voluntary reporting, and facilitating the aggregation of large volumes of data which in turn aids in identifying patterns of patient safety events. Under the Patient Safety Act, the imposition of civil monetary penalties is authorized for breaches of its confidentiality provisions. The confidentiality protections of patient safety information are to be implemented in a way that does not interfere with other health care reporting obligations of providers, e.g., under State or local laws.

Dated: February 13, 2008.

**Carolyn M. Clancy,**  
*AHRQ, Director.*

[FR Doc. 08-776 Filed 2-20-08; 8:45am]

BILLING CODE 4160-90-M

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Administration for Children and Families

#### Proposed Information Collection Activity; Comment Request

*Proposed Project:*

*Title:* Exploration of Low-Income Couples' Decision-Making Processes.  
*OMB No.:* New Collection.

*Description:* The Administration for Children and Families (ACF), U.S. Department of Health and Human Services (HHS), is proposing a data collection activity as part of the Exploration of Low-Income Couples' Decision-Making (CDM) Processes study. This project will gather important

information that will be useful for improving social services delivery approaches for working with individuals in couple relationships. The proposed collection will consist of two elements: (1) Focus groups with low-income couples; and (2) a telephone survey and observation of low-income

couples. These data collection efforts will examine sources of conflict and assess decision-making processes among low-income couples—especially in relation to issues directly addressed by social service programs (e.g., employment, housing).

*Respondents:* Low-income couples.

ANNUAL BURDEN ESTIMATES

Instrument	Annual number of respondents	Number of responses per respondent	Average burden hours per response	Estimated annual burden hours
Focus Group Discussion .....	16	1	2	32
Telephone Survey .....	80	1	.333333	27
Home Visit Setup and Administration of Oral History Interview and Decision Payoff Ratings .....	80	1	.666666	53
Paper Tower Task .....	80	1	.5	40
Economic Decision Task—Revealed Differences .....	80	1	.25	20
Interpersonal Conflict Discussion .....	80	1	.25	20
Video Recall Task .....	80	1	.83	66

Estimated Total Annual Burden Hours: 258.

In compliance with the requirements of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded by writing to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. E-mail address: [infocollection@acf.hhs.gov](mailto:infocollection@acf.hhs.gov). All requests should be identified by the title of the information collection.

The Department specifically requests comments 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) 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. Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Dated: February 12, 2008.

**Brendan C. Kelly,**

*Reports Clearance Officer.*

[FR Doc. 08-777 Filed 2-20-08; 8:45 am]

**BILLING CODE 4184-01-M**

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

[Docket No. FDA-2008-N-0082]

**Animal Drug User Fee Act; Public Meeting; Request for Comments**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice of public meeting; request for comments.

**SUMMARY:** The Food and Drug Administration (FDA, the agency) is publishing proposed recommendations for the reauthorization of the Animal Drug User Fee Act of 2003 (ADUFA) for fiscal years (FY) 2009 to 2013. These proposed recommendations were developed after a public meeting with stakeholders and discussions with regulated industry. ADUFA, enacted November 18, 2003, directs FDA to publish these proposed recommendations in the **Federal Register**; hold a meeting at which the public may present its views on such recommendations; and provide a period of 30 days for the public to provide written comments on such recommendations.

*Dates and Time:* The public meeting will be held on March 11, 2008, from 1 p.m. to 3:30 p.m.

*Location:* The public meeting will be held at 7519 Standish Pl., third floor,

rm. A, Rockville, MD 20855. There is parking near the building. Photo identification is required to clear building security.

*Contact Person:* Roxanne Schweitzer, Center for Veterinary Medicine (HFV-10), Food and Drug Administration, 7529 Standish Pl., Rockville, MD 20855, 240-276-9705, FAX: 240-276-9744, e-mail: [Roxanne.Schweitzer@fda.hhs.gov](mailto:Roxanne.Schweitzer@fda.hhs.gov).

*Registration:* To ensure there is sufficient room we ask that you pre-register. Furthermore, to assist us in scheduling, we ask that you notify us through the registration process if you wish to make a public comment at the meeting. To register, please send an electronic mail message to [roxanne.schweitzer@fda.hhs.gov](mailto:roxanne.schweitzer@fda.hhs.gov) by March 4, 2008. Your e-mail should include the following information: Name, Company, Company Address, Company Telephone Number, and E-mail Address. You will receive a confirmation within 2 business days.

FDA also will accept walk-in registration at the meeting site, but space is limited, and the agency will close registration when maximum seating capacity (approximately 500) is reached. FDA will try to accommodate all persons who wish to make a public comment at the meeting, including those who register at the meeting site, however, the time allotted for public comments may depend on the number of persons who wish to speak.

Additionally, please notify FDA (see *Contact Person*) if you need any special accommodations (such as wheelchair access or a sign language interpreter) at least 7 days in advance.

*Comments:* To ensure consideration of your comments regarding these proposed recommendations, you should