1. Samuel T. Sicard, individually and as trustee of the Samuel M. Sicard Living Trust, Fort Smith, Arkansas; to retain ownership of First Bank Corp., and thereby indirectly retain ownership of The First National Bank of Fort Smith, both in Fort Smith, Arkansas.

Board of Governors of the Federal Reserve System, November 17, 2011.

#### Robert deV. Frierson,

Deputy Secretary of the Board. [FR Doc. 2011–30106 Filed 11–21–11; 8:45 am]

BILLING CODE 6210-01-P

#### FEDERAL RESERVE SYSTEM

### Formations of, Acquisitions by, and Mergers of Bank Holding Companies

The companies listed in this notice have applied to the Board for approval, pursuant to the Bank Holding Company Act of 1956 (12 U.S.C. 1841 et seq.) (BHC Act), Regulation Y (12 CFR Part 225), and all other applicable statutes and regulations to become a bank holding company and/or to acquire the assets or the ownership of, control of, or the power to vote shares of a bank or bank holding company and all of the banks and nonbanking companies owned by the bank holding company, including the companies listed below.

The applications listed below, as well as other related filings required by the Board, are available for immediate inspection at the Federal Reserve Bank indicated. The application 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.

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 December 16, 2011.

A. Federal Reserve Bank of San Francisco (Kenneth Binning, Vice President, Applications and Enforcement) 101 Market Street, San Francisco, California 94105–1579:

1. American Start-Up Financial Institutions Investments, I, L.P., and CKH Capital, Inc., both in Monterey Park, California; to become bank holding companies by acquiring up to 62 percent of the voting shares of New Omni Bank, National Association, Alhambra, California.

In connection with this application, Applicants also have applied to retain 5.9 percent interest of the voting shares of First PacTrust Bancorp, Inc., and thereby indirectly retain Pacific Trust Bank, both in Chula Vista, California, and engage in operating as savings and loan association, pursuant to section 225.28(b)(4)(ii) of Regulation Y.

Board of Governors of the Federal Reserve System, November 17, 2011.

#### Robert deV. Frierson,

Deputy Secretary of the Board.
[FR Doc. 2011–30105 Filed 11–21–11; 8:45 am]
BILLING CODE 6210–01–P

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Centers for Disease Control and Prevention

[60Day-12-12AM]

## 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 (404) 639-5960 and send comments to Daniel Holcomb, CDC Reports Clearance Officer, 1600 Clifton Road, MS-D74, Atlanta, GA 30333 or send an email to omb@cdc.gov.

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. Written comments should be received within 60 days of this notice.

#### **Proposed Project**

Prospective Birth Cohort Study Involving Environmental Uranium Exposure in the Navajo Nation (U01)— New—National Center for Environmental Health (NCEH) and Agency for Toxic Substances and Disease Registry (ATSDR), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The Navajo Nation includes 16 million acres of New Mexico, Utah and Arizona. It is the largest Alaska Native/ American Indian Reservation in the United States. From 1948 to 1986, many uranium mining and milling operations took place in the Navajo Nation, leaving a large amount of uranium contamination on the reservation. Several studies have reported that uranium mostly damages the kidneys and urinary system. However, there is not much research data on uranium exposure and poor birth and reproductive health outcomes. Research involving prenatal exposure to uranium may help to understand and prevent some unfavorable child and maternal health outcomes.

There are important health differences concerning birth outcomes and prenatal care in the Navajo Nation. According to the Indian Health Service Regional Differences in Indian Health 2002-2003 Edition, the infant death rate among the Navajo people is 8.5 deaths per 1000 live births, compared to 6.9 deaths per 1000 live births among all races. Only 61% of Navajo mothers with live births received prenatal care in the first trimester as compared to 83% of all U.S. mothers. Early and regular prenatal care is a major predicator of positive birth outcomes. Due to the health differences in birth outcomes and the chance for environmental uranium exposure in the Navajo Nation, ATSDR decided that the upcoming study must include education of women and their families about the importance of prenatal care and the potential poor health risks associated with exposure to uranium.

The House Committee on Oversight and Government Reform requested that federal agencies develop a plan to address health and environmental impacts of uranium contamination in the Navajo Nation. As a result of this request, ATSDR awarded a research cooperative agreement to University of New Mexico Community Environmental Health Program (UNM-CEHP) entitled "A Prospective Birth Cohort Study Involving Environmental Uranium Exposure in the Navajo Nation (U01)," in August 2010. ATSDR and UNM-CEHP are working with the Navajo Area Indian Health Service (NAIHS), Navajo Nation Division of Health (NNDOH), Navajo Nation Environmental Protection Agency (NNEPA), and Navajo culture

and language specialists to carry out the study. The study will examine reproductive outcomes in pregnant women, follow and assess their children from birth to 1 year of age, and create a system to follow up the infants through childhood up to 6 years of age to evaluate the impact of uranium exposure on biological and psychosocial endpoints. Biological sample analysis, surveys, and developmental screenings will be performed during this research period for each participant.

In addition to investigating the role of uranium and other chemicals in the environment on birth outcomes and development, the prospective study may aid in understanding causes and prevention measures of chronic conditions. Several research studies have shown that exposure to chemicals in the environment during prenatal and postnatal periods can affect the development of adult chronic diseases. The study will also provide broad public health benefits for Navajo communities through outreach and education on environmental prenatal risks and early assessment. Referrals will also be provided for known developmental delays.

Participants will include Native American mothers from age 14 to 45 with verification of pregnancy who have lived in the study area for at least 5 years. Also, participants must consent to receive prenatal care and deliver at one of the healthcare facilities that are taking part in the study (Northern Navajo Medical Center, Chinle Comprehensive Health Care Facility, Gallup Indian Medical Center, Tuba City Regional Health-Care Corporation, or Tséhootsooí Medical Center). Fathers will be included in the study with consent regardless of age or residence. We estimate that 550 pregnant women and fathers per year must be enrolled in the study to obtain adequate statistical power. A 10% pregnancy loss will be assumed, which would result in 500 live births per year. Therefore, the total anticipated sample size is 1,500 motherinfant pairs over the three years of the

The survey instruments for pregnant mothers include the following: Enrollment Survey, Nutritional Assessment/Food Intake Questionnaire, Ages and Stages Questionnaire (ASQ–I), Mullen Stages of Early Development (MSEL), and Postpartum Surveys. An

enrollment survey for fathers who agree to participate will also be administered. Community Health and Environmental Research Specialists (CHERS) will administer surveys using a CDCapproved electronic data entry system. Survey instruments were designed to collect demographic information, assess potential environmental health risks, and mother-child interactions. The survey instruments were developed based on previous surveys conducted by Dine' Network for Environmental Health (DiNEH) Project, the National Children's Study, and by other birth cohort studies that have been conducted among other indigenous populations. The final format of the survey instruments was modified based on review and input from the Navajo Nation community liaison group and associated Navajo staff to address issues such as cultural sensitivity, comprehension and language translation.

There is no cost to the respondents other than their time to participate in the study. The total estimated annual burden hours equals 3550.

Estimated Annualized Burden Hours

Type of respondent	Form name	Number of re- spondents	Number of responses per respondent	Average bur- den response (hours)	Total burden (hours)
Mother	Enrollment Survey	550	1	2	1100
	Ages and Stages Questionnaire (2,6,9 12 months).	500	4	15/60	500
	Mullen Stages of Early Development	500	1	15/60	125
	Postpartum Survey (0 months)	500	1	1	500
	Post-partum Survey (2, 6, 9, 12 months).	500	4	15/60	500
Father	Enrollment Survey	550	1	90/60	825
Total					3550

Dated: November 16, 2011.

#### Daniel Holcomb,

Reports Clearance Officer, Centers for Disease Control and Prevention.

[FR Doc. 2011–30103 Filed 11–21–11; 8:45 am] BILLING CODE 4163–18–P

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### **National Institutes of Health**

### Government-Owned Inventions; Availability for Licensing

**AGENCY:** National Institutes of Health, Public Health Service, HHS.

**ACTION:** Notice.

**SUMMARY:** The inventions listed below are owned by an agency of the U.S. Government and are available for

licensing in the U.S. in accordance with 35 U.S.C. 207 to achieve expeditious commercialization of results of federally-funded research and development. Foreign patent applications are filed on selected inventions to extend market coverage for companies and may also be available for licensing.

ADDRESSES: Licensing information and copies of the U.S. patent applications listed below may be obtained by writing to the indicated licensing contact at the Office of Technology Transfer, National Institutes of Health, 6011 Executive Boulevard, Suite 325, Rockville, Maryland 20852–3804; telephone: (301) 496–7057; fax: (301) 402–0220. A signed Confidential Disclosure Agreement will be required to receive copies of the patent applications.

# Medical Device for Intraocular Injection of Therapeutics and Fluid Sampling

Description of Technology: The National Institutes of Health seeks research collaboration and commercialization partners for a medical device for administering therapeutics into the eye to treat a variety of ocular diseases including diabetic retinopathy, retinal vein occlusion, and macular degeneration. The device is a dual function needle that can both inject and sample ocular fluid at the same injection site. The needle includes a hub portion in communication with a needle portion through a lumen that may be used as a conduit to inject a therapeutic into an injection site. A sample chamber, with an optional absorbent material, is